WO 01/05574 PCT/US00/19320

achieved over the tested range is called the ultimate hot tack strength. the point at which the hot tack strength achieves 2 N/2.54cm. The ultimate strength 10° Celsius increments from 80-150°C Celsius. Hot tack initiation temperature is defined as

ultimate hot tack (4.26 for 7B, 2.78 for 7A). The ultimate hot tack strength is found to that is decrease in hot tack initiation temperature of 12°C (83°C for 7B, 95°C for 7A) and in increase with increasing fringe heights. Example 6. As compared to the base film 7A, the fringed film 7B shows an improvement, height (H) is determined via optical microscopy according to the method described in and have the structure and composition of film 1A described in Example 1, above. Fringe 250 microns (7B) are tested for their hot tack performance. Both films are three-layer films A smooth base film (7A) (control) and a fringed film (7B) having a fringe height (H) of

Example 8: Water Repellency of Fringed Film

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water to bead up on the surface. The water contact angle is measured using pH 7 buffer the fringed surface microstructure has on the water repellency or the ability to enable the related to a wetting tension via a conversion chart. The test is used to compare the effect which the side of the water droplet contacts the substrate. Water contact angle can then be directly with a protractor by using a tangential alignment of a cross-hair line to the point at sample, and the contact angle values are then averaged. The contact angle is measured method 1 in 3.1.2.1. In this test method water droplets are placed on the surface of a film the water contact angle test is utilized. Testing is done according to ASTM-D5946 utilizing To examine the water repellency or water management characteristics of fringed film

angle, indicating an increased ability to bead up water on the surface. Such effect contact angle determined for film 8A is 137.3, the average water contact angle for film 8B is characteristics as films 6B and 6C described in Example 6 and are referred to as film 8A important for applications requiring water repellency, such as raincoats and other water 155.4. The results show that increased fringe length results in an increased water contact Two separate areas per film were evaluated and the values averaged. The average water (fringe height (H) of 70 microns) and 8B (fringe height (H) of 95 microns) in this example. gloves (prevent formation of a slippery surface). repellant outerwear, house wraps, feminine hygiene and diaper top sheets, and (latex) The test is performed on films having the same structure, composition and fringe

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WHAT IS CLAIMED IS:

displays a surface microstructure, which layer is a thermoplastic polymeric material and least partially hollow, in a density of 1000 or more per square centimeter. characterized by fringes, said fringes being non-perforated crater-like peaks which are at A mono-layer or multi-layer film, sheet, or coating wherein at least one layer

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(H) in the range of from 40 microns to 1 millimeter. A film, sheet or coating according to claim 1, wherein the fringes have a height

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- depth ratio (•) of 1.3 or lower, as determined by optical surface profilometry. A film, sheet or coating according to claim 1, wherein the fringes have a hollow
- diameter ratio (*) of 1 or lower, as determined by optical surface profilometry A film, sheet or coating according to claim 1, wherein the fringes have a hollow
- Hollowness Index (•) of 100 or lower, as determined by optical surface profilometry. A film, sheet or coating according to claim 1, wherein the fringes have
- aspect ratio (A) of between 1 and 5. The film, sheet or coating according to claim 1, wherein the fringes have an
- material is cured, irradiated or cross-linked. 7. The film, sheet, or coating according to claim 1, wherein the thermoplastic

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- surface microstructure on both sides 8. The film, sheet, or coating according to claim 1, wherein the layer displays a
- or coating. 9. The film, sheet, or coating according to claim 1, which is a mono-layer film, sheet

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coating. 10. The film, sheet, or coating according to claim 1, which is a multi-layer film, sheet or

- microstructure is on an outer layer. 11. The film, sheet, or coating according to claim 10, wherein the surface
- coating, wherein the surface-structured layer is an interlayer. The film, sheet, or coating according to claim 1 which is a multi-layer film, sheet or
- biaxially oriented polypropylene film. layers is a fringed layer and at least one of the inner layers is an oriented film, preferably a 13. The film, sheet or coating according to claim 10, wherein at least on of the outer

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- Ö a foamed layer. The film, sheet or coating according to claim 13, wherein at least one of the layers
- 15 elastic The film, sheet, or coating according to claim 1, wherein at least one layer is
- The film, sheet, or coating according to claim 1, which is oriented
- permeable and liquid impermeable 17. The film, sheet, or coating according to claim 1, wherein at least one layer is vapor

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- 18. The film, sheet, or coating according to claim 1 which is printed or imprinted
- 2 5 with an abrading device, corona treatment, curing, irradiation and crosslinking has been subjected to a post treatment step selected from the group consisting of treatment 19. The film, sheet or coating according to claim 1 wherein the surface microstructure
- A composite comprising the film, sheet or coating according to claim 1.
- 21. The composite according to claim 20 which is a laminate

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coating according to claim 1. 22. An article of manufacture comprising a mono-layer or multi-layer film, sheet, or

- 23. The article of manufacture according to claim 22, which is a glove
- The article of manufacture according to claim 22, which is a hygienic product.
- bag, preferably an ostomy bag. 25. The article of manufacture according to claim 22, which is a medicinal collection
- product. The article of manufacture according to claim 22, which is a floor or wall covering

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- touch. The article of manufacture according to claim 22 which has a soft, textile-like
- The article of manufacture according to claim 22 which is water repellant.

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- properties 29. The article article of manufacture according to claim 28 which has anti-skid
- 20 capturing or storing properties 30. The article of manufacture according to claim 22 which has enhanced carrying,
- 31. The article of manufacture according to claim 22 which is heat resistant.
- 25 comprising 32. An process for making a film, sheet, or coating according to claim 1, said process
- providing a precursor film, sheet, or coating with a surface characterized by a pattern of peaks and valleys, and
- subjecting said precursor to mechanical treatment under conditions allowing the formation of a fringed surface microstructure.

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formed in a continuous compression molding process and the mechanical treatment 33. The process according to claim 32, wherein the precursor film, sheet, or coating is

the film, sheet, or coating from a matrix surface. comprises application of a tractive force to said precursor which is applied during release of

- between 20 and 170 degrees relative to the matrix surface at or below the Vicat softening point of the thermoplastic material and at a release angle of comprises peeling the film, sheet, or coating off the matrix surface at a temperature which is The process according to claim 32, wherein application of the tractive
- 35. The process according to claim 32, wherein the surface is a foam

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- calander, to make the film, sheet or coating according to claim 1. 36. A device comprising a matrix surface and a counter surface, preferably a
- manufacture having a soft, textile-like touch. 37. Use of the film, sheet or coating according to claim 1 to make an article of

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- 38. Use of the film, sheet or coating according to claim 1 in automotive applications.
- water repellant. 39. Use of the film, sheet or coating according to claim 1 to make an article which is

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- 40. Use of the film, sheet or coating according to claim 1 for packaging.
- carrying, capturing or storing properties. 41. Use of the film, sheet or coating according to claim 1 in an article with enhanced

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- anti-skid properties 42. Use of the film, sheet or coating according to claim 1 to make an article which has
- enhanced heat resistance 43. Use of the film, sheet or coating according to claim 1 to make an article which has

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ď matte surface appearance 44. Use of the film, sheet or coating according to claim 1 to make an article which has

- 45. Use of the film, sheet or coating according to claim 1 as a filtration medium.
- claim 1. 46. Use of a thermoplastic material to make the film, sheet or coating according to

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preferably poly(lactide). 47. Use of a thermoplastic material according to claim 46 which is biodegradable,

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(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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16 July 1999 (16.07.1999) 14 September 1999 (14.09.1999)

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International Application Number:

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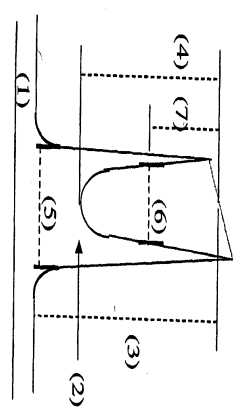
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(81) AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, II, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, Designated States (national): AE, AG, AL, AM, AT, AU, ug, us, uz, yu, za, zw.

84 Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IF, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

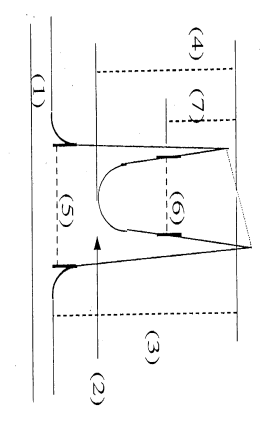
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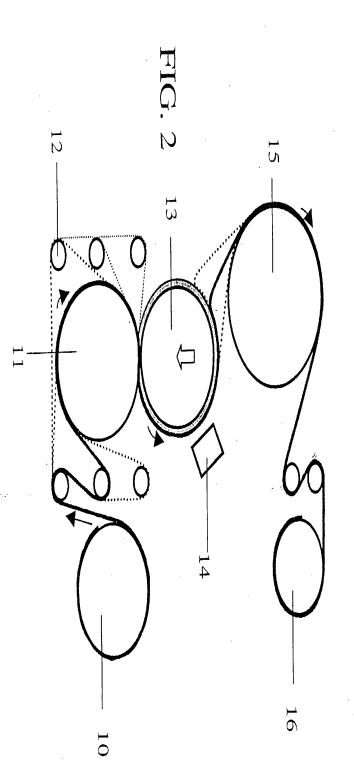
(54) Title: FRINGED SURFACE STRUCTURES OBTAINABLE IN A COMPRESSION MOLDING PROCESS



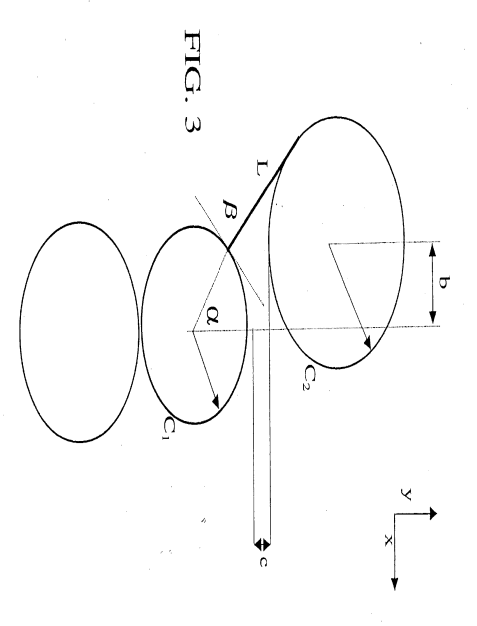
sheet, or coating articles items and uses thereof. microstructure at least one layer multi-layer films, sheets, (57) Abstract: of manufacture Ω, Disclosed are mono-layer a process for making these comprises a fringed surface comprising Further disclosed Ŷ coatings wherein

PCT/US00/19320





PCT/US00/19320



DECLARATION FOR PATENT APPLICATION

As an undersigned inventor, I hereby declare that:

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	[x]	I believe (check one) []
original and first inventors	I am a joint inventor and the below named inventors are the	I am the original, first and sole inventor

of the subject matter which is claimed and for which a patent is sought on the invention entitled

FRINGED SURFACE STRUCTURES OBTAINABLE IN A COMPRESSION MOLDING PROCESS

and was amended on(if applicable)	the specification of which (check one) []	/hich [X]	is attached hereto. was filed on <u>January 15, 2002</u> , as Application Serial No. 10/031,304,
and was amended on (if applicable)		[x]	was filed on <u>January 15, 2002</u> , as Application Serial No. 10/031 304
			_1

specification, including the claims, as amended by any amendment referred to above. I further declare that I have reviewed and understand the contents of the above identified

which is claimed as defined in 37 C.F.R. §1.56. "the Office") all information known to me to be material to patentability of the subject matter I acknowledge the duty to disclose to the United States Patent and Trademark Office (hereinafter

on which priority is claimed: application for patent or inventor's certificate having a filing date before that of the application patent or inventor's certificate indicated below and have also identified below any foreign I hereby claim foreign priority benefits under 35 U.S.C. § 119 of any foreign application(s) for

Prior Foreign Application(s) Country Day/Month/Year Filed

Priority
Claimed
Yes No

I hereby claim the benefit under 35 U.S. C. §119(e) of any United States provisional application listed below:

Provisional Application Serial No. 60/144,306 – 60/153,793 –

The state of the s

Filing Date
July 16, 1999
September 14, 1999

acknowledge the duty to disclose to the Office all information known to me to be material to patentability as defined in 37 C.F.R. § 1.56, which became available between the filing date of the prior application and the national or PCT international filing date of this application: international application in the manner provided by the first paragraph of 35 U.S.C. § 112, I matter of each of the claims of this application is not disclosed in the prior United States or PCT international application designating the United States listed below and, insofar as the subject I hereby claim the benefit under 35 U.S.C. § 120 of any United States application(s) and PCT

Application Serial No. PCT/US00/19320

Filing Date
July 17, 2000 ~

Status (patented, pending, abandoned) PCT

Address all correspondence and telephone calls to the following:

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statements made on information and belief are believed to be true; and further, that these statements were made with the knowledge that willful false statements and the like are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and may jeopardize the validity of the application or any patent issued thereon. I hereby declare that all statements made herein of my own knowledge are true and that all

Post Office Address: Same (enter "same" if mailing address is same as residence address)	Citizenship: AUSTRIA —	Residence: Felsenrainstrasse 7a, CH-8832 Wollerau SZ, Switzerland C++×.	Inventor's Signature 14/6/2002 Date	Thomas T. Allgeuer
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Inventor's Signature Adams Drive Howsen Tyle Togs Citizenship: TURKEY Post Office Address: Same (enter "same" if mailing address is same as residence address)	
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ddress)		X	Date	

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Post Office Address: (ent	ip: 45.	e: 1506 Ft	Inventor's Signature	-OFull name of seventh joint inventor	
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8-60 Full name of eighth joint inventor: Werney Wagner Inventor's Signature Dy benus tosapues

July 22, 2002 Date

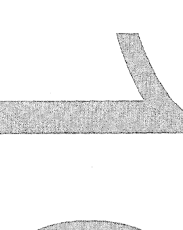
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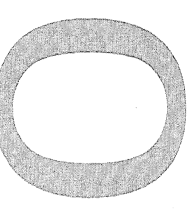
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Level - 2 Version 1.1 Updated - 8/01/01

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371 Application As-Filed

Level - 1 Version 1.1 Updated - 8/01/01



IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Jacquelyne A.; KELCH, Robert H.; LIANG, Wenbin; TORRES, Antonio; WAGNER, Werner ALLGEUER, Thomas T.; BENSASON, Selim; CHUM, Pak-Wing S.; DEGROOT,

Filed: Concurrently Herewith

FRINGED SURFACE STRUCTURES OBTAINABLE IN A COMPRESSION MOLDING PROCESS

International Application No.: PCT/US00/19320

International Filing Date: July 17, 2000

Priority Date Claimed: July 16, 1999

Serial No.: Unknown

Group Art Unit: Unknown

Examiner: Unknown

Atty Dkt: ADVA:005

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Sign	1	indigated below and is address/6:) Assistant Commissioner of Patents, Washington, D.C. 20231.	Postal Service "EXPRESS MAIL POST OFFICE TO ADDRESSEE" service, postage prepaid, under 37 CFR 1.10 on the date	EXPRESS MAIL NUMBER: EV 044385928 US I hereby certify that this paper or fee is being
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Assistant Commissioner for Patents Washington, DC 20231

Dear Sir:

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the

following items and other information:

- Mark This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.
- 'n This express request to begin national examination procedures (35 U.S.C. 371(f) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and

PCT Articles 22 and 39(1).

☐ This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371.

- 4 A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.
- Ġ A copy of the International Application as filed (35 U.S.C. 371(c)(2))

531 Rac'd PCT/P 15 JAN 2002

SSI HEC OTENT IS JAN 200
 a. is transmitted herewith (required only if not transmitted by the International Burcau). b. has been transmitted by the International Bureau. c. is not required, as the application was filed in the United States Receiving Office (RO/US); however, a copy is submitted herewith, including rectified FIGS. 1-3.
6. A translation of the International Application into English (35 U.S.C. 317(c)(2)).
 7. Amendments to claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)). a. are transmitted herewith (required only if not transmitted by the International Bureau). b. have been transmitted by the International Bureau. c. have not been made; however, the time limit for making such amendments has NOT expired. d. have not been made and will not be made.
8. A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
9. An oath or declaration of the inventor(s) 35 U.S.C. 371(c)(4)).
10. [] a translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 317(c)(5)).
Items 11 to 16 below concern document(s) or information included:
11. An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
12. An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
13. A FIRST preliminary amendment. Please enter this amendment prior to calculating the file fee.
☐ A SECOND or SUBSEQUENT preliminary amendment.
14. A substitute specification.
15. A change of power of attorney and/or address letter.
16. Other items or information: Request for Priority
17. The following fees are submitted:

537 Rec'd PC1/F 15 JAN 2002

\$ 710.00 \$1000.00 \$ 100.00 \$ 860.00	(§1.492(a)(2)) International preliminary examination fee not paid to USPTO, but an international search fee was paid to the USPTO
\$ 690.00	(§1.492(a)(1)) International preliminary examination fee paid to USPTO
	BASIC NATIONAL FEE (37 CFR 1.492(a) (1) – (5)):

FEE CALCULATION:

CLAIMS	(1) FOR	(2) NUMBER FILED	(3) NUMBER EXTRA	(4) RATE	(5) CALCULATIONS
	Total Claims (37 CFR 1.16(e))	47 - 20 =	27	x \$18.00	\$ 486.00
	Independent Claims (37 CFR 1.16(b))	_3 3 =	0	× \$80.00	\$0
	MULTIPLE DEPENDENT CLAIMS (if applicable) (37 CFR 1.16(d))	S (if applicable) (37 CFR	1.16(d))	x \$ 270.00	\$0
				Basic Fee (from above)	\$ 860.00
			Total of above Calculations	ulations =	\$1346.00.00
	Reduction by 50% for filing by Small Entity (Note: 37 CFR 1.9, 1.27, 1.28)	tity (Note: 37 CFR 1.9, 1.	27, 1.28)		\$0
	Processing fee of \$130.00 for furnishing the English translation later than \square 29 \square 30 months claimed priority date (37 CFR 1.492(f)).	the English translation late	r than □ 29 □30 r	nonths from the earliest	\$0
	Surcharge of \$130.00 for furnishing the oath or declaration later than \(\subseteq 20 \) \(\subseteq 30 \) months from claimed priority date (37 CFR 1.492(e)).	ath or declaration later tha	ın 🗌 20 🔲 30 mon	ths from the earliest	\$0
		TO	TOTAL NATIONAL	VAL FEE =	49
	Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31) \$40.00 per property.	at (37 CFR 1.21(h)). The a	assignment must be a	ccompanied by an	49
		ТС	TOTAL FEES ENC	ENCLOSED =	\$1346.00

- a. A check in the amount of $\underline{\$1346.00}$ to cover the above fees is enclosed
- <u></u> ☐ If the check is inadvertently omitted, or should any additional fees under 37 CFR §1.16 to 1.21, overpayment be included herein, the Commissioner is authorized to deduct or credit said fees from or to Deposit Account No. 10-1205. §1.492, or any other rule, be required for any reason relating to the enclosed materials, or should an
- X incorporating a petition for extension of time for the appropriate length of time, and the In accordance with 37 CFR 1.136(a)(3), the Commissioner is authorized to treat any concurrent or Commissioner is authorized to deduct any requisite fees under 37 CFR 1.16 to 1.21 from Deposit Account No. 10-1205. future reply that requires a petition for an extension of time under 37 CFR 1.126(a) to be timely, as

Please forward all correspondence to the undersigned at the address below. This application is submitted prior to the 30-month deadline in the international PCT application.

Respectfully submitted,

Robert M. O'Keefe Reg. No. 35,630

Attorney for Applicants

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: ALLGEUER, Thomas T.; BENSASON, Selim; CHUM, Pak-Wing S.; DEGROOT, Jacquelyne A.; KELCH, Robert H.; LIANG, Wenbin; TORRES, Antonio; WAGNER, Werner

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FRINGED SURFACE STRUCTURES OBTAINABLE IN A COMPRESSION

MOLDING PROCESS

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Assistant Commissioner For Patents Washington, D.C. 20231

Dear Sir:

PRELIMINARY AMENDMENT

dependent claims thereby correcting matters of form. Please enter this amendment prior to calculating the filing fee due. Upon entry the amendment, the pending claims shall constitute claims 1-47. This paper is submitted amend the claims to remove move certain recitations into new

In the specification:

On page 1, please enter the following new paragraph immediately after the title of the

September 14, 1999, incorporated herein by reference. 60/144,306 filed July 16, 1999 and to US provisional application serial number 60/153,793, filed filed July 17, 2000, which claims priority to US provisional application serial number (New) This application is a 35 U.S.C. §371 nationalized application of PCT/US00/19320,

In the Claims:

II of the PCT. by Rule 121(c). The amendments are of the claims submitted on August 6, 2001, during Chapter providing an indication of the changes relative to the prior version of the claims, as now required the pending claims are provided below. Attached at the end of this paper is an Appendix Please enter the following amended and new claims. The rewritten clean versions of all

- being empty and have a height of at least about 40 microns or more, in a density of 1000 or are at least partially hollow with at least about 25 percent of the volume at the top of the peak fringes, said fringes being characterized in that they are non-perforated crater-like peaks which a surface microstructure, which layer is a thermoplastic polymeric material and characterized by more per square centimeter. 1. A mono-layer or multi-layer film, sheet, or coating wherein at least one layer displays
- range of from 40 microns to 1 millimeter. A film, sheet or coating according to claim 1, wherein the fringes have a height in the
- of 1.3 or lower, as determined by optical surface profilometry. ratio, which is the ratio of the average inner height to the average maximum height of the fringe 3. A film, sheet or coating according to claim 1, wherein the fringes have a hollow depth

- diameter ratio, which is ratio of the diameter of the hollow center at half height and the diameter at the bottom of the fringe, of 1 or lower, as determined by optical surface profilometry. 4. A film, sheet or coating according to claim 1, wherein the fringes have a hollow
- Index, which is calculated by multiplying the hollow depth ratio with the hollow diameter ratio, of 100 or lower, as determined by optical surface profilometry. 5. A film, sheet or coating according to claim 1, wherein the fringes have a Hollowness
- ratio, which is the ratio of the fringe height and the fringe diameter, of between 1 and 5. 6. The film, sheet or coating according to claim 1, wherein the fringes have an aspect
- cured, irradiated or cross-linked. 7. The film, sheet, or coating according to claim 1, wherein the thermoplastic material is
- microstructure on both sides. The film, sheet, or coating according to claim 1, wherein the layer displays a surface
- 9. The film, sheet, or coating according to claim 1, which is a mono-layer film, sheet or
- 10. The film, sheet, or coating according to claim 1, which is a multi-layer film, sheet or
- is on an outer layer. 11. The film, sheet, or coating according to claim 10, wherein the surface microstructure
- coating, wherein the surface-structured layer is an interlayer. 12. The film, sheet, or coating according to claim 1 which is a multi-layer film, sheet or
- biaxially oriented polypropylene film. layers is a fringed layer and at least one of the inner layers is an oriented film, preferably a The film, sheet or coating according to claim 10, wherein at least on of the outer

- 14. The film, sheet or coating according to claim 13, wherein at least one of the layers is a
- 15. The film, sheet, or coating according to claim 1, wherein at least one layer is elastic.
- 16. The film, sheet, or coating according to claim 1, which is oriented
- permeable and liquid impermeable. 17. The film, sheet, or coating according to claim 1, wherein at least one layer is vapor
- 18. The film, sheet, or coating according to claim 1 which is printed or imprinted
- abrading device, corona treatment, curing, irradiation and crosslinking. been subjected to a post treatment step selected from the group consisting of treatment with an 19. The film, sheet or coating according to claim 1 wherein the surface microstructure has
- more, in a density of 1000 or more per square centimeter. the volume at the top of the peak being empty and have a height of at least about 40 microns or perforated crater-like peaks which are at least partially hollow with at least about 25 percent of material and characterized by fringes, said fringes being characterized in that they are nonleast one layer displays a surface microstructure, which layer is a thermoplastic polymeric 20. A composite comprising a mono-layer or multi-layer film, sheet, or coating wherein at
- 21. The composite according to claim 20 which is a laminate.
- about 40 microns or more, in a density of 1000 or more per square centimeter. about 25 percent of the volume at the top of the peak being empty and have a height of at least in that they are non-perforated crater-like peaks which are at least partially hollow with at least thermoplastic polymeric material and characterized by fringes, said fringes being characterized coating wherein at least one layer displays a surface microstructure, which layer is a 22. An article of manufacture comprising a mono-layer or multi-layer film, sheet, or

- 23. The article of manufacture according to claim 22, which is a glove
- 24. The article of manufacture according to claim 22, which is a hygienic product.
- collection bag[, preferably an ostomy bag]. 25. (Amended) The article of manufacture according to claim 22, which is a medicinal
- product. 26. The article of manufacture according to claim 22, which is a floor or wall covering
- 27. The article of manufacture according to claim 22 which has a soft, textile-like touch.
- 28. The article of manufacture according to claim 22 which is water repellant.
- 29. The article article of manufacture according to claim 28 which has anti-skid properties
- capturing or storing properties. 30. The article of manufacture according to claim 22 which has enhanced carrying,
- 31. The article of manufacture according to claim 22 which is heat resistant.
- claim 1, said process comprising 32. A process for making the mono-layer or multi-layer film, sheet, or coating according to
- of peaks and valleys in a continuous compression molding process, and forming a precursor film, sheet, or coating with a surface characterized by a pattern
- subjecting said precursor to mechanical treatment comprising the application of a matrix surface under conditions allowing the formation of a fringed surface microstructure. tractive force which is applied during release of the film, sheet or coating from a

- 170 degrees relative to the matrix surface. the Vicat softening point of the thermoplastic material and at a release angle of between 20 and peeling the film, sheet, or coating off the matrix surface at a temperature which is at or below 33. The process according to claim 32, wherein application of the tractive force comprises
- 34. The process according to claim 32, wherein the precursor is a foam.
- manufacture having a soft, textile-like touch. 35. Use of the film, sheet or coating according to claim 1 to make an article of
- 36. Use of the film, sheet or coating according to claim 1 in automotive applications.
- repellant. 37. Use of the film, sheet or coating according to claim 1 to make an article which is water
- 38. Use of the film, sheet or coating according to claim 1 for packaging.
- carrying, capturing or storing properties. 39. Use of the film, sheet or coating according to claim 1 in an article with enhanced
- anti-skid properties. 40. Use of the film, sheet or coating according to claim 1 to make an article which has
- enhanced heat resistance 41. Use of the film, sheet or coating according to claim 1 to make an article which has
- matte surface appearance. 42. Use of the film, sheet or coating according to claim 1 to make an article which has a
- 43. Use of the film, sheet or coating according to claim 1 as a filtration medium.

- 44. Use of a thermoplastic material to make the film, sheet or coating according to claim
- biodegradable. 45. (Amended) Use of a thermoplastic material according to claim 44 which is
- 46. (New) The article of manufacture according to claim 22, which is an ostomy bag.
- material is a poly(lactide). 47. (New) Use of a thermoplastic material according to claim 44 which themroplastic

CONCLUSION

and compact prosecution of the application. number indicated below with any questions or comments, or to otherwise facilitate expeditious No. 10-1205/ADVA:005. The examiner is invited to contact the undersigned at the phone enclosed materials, the Commissioner is authorized to deduct such fees from Deposit Account Should any fees under 37 CFR 1.16-1.21 be required for any reason relating to the

Respectfully submitted,

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531 Rec'd PCI/PI 15 JAN 2002

APPENDIX MARKED UP VERSION OF AMENDMENTS AS REQUIRED BY RULE 121

- more per square centimeter. being empty and have a height of at least about 40 microns or more, in a density of 1000 or are at least partially hollow with at least about 25 percent of the volume at the top of the peak fringes, said fringes being characterized in that they are non-perforated crater-like peaks which a surface microstructure, which layer is a thermoplastic polymeric material and characterized by 1. A mono-layer or multi-layer film, sheet, or coating wherein at least one layer displays
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- at the bottom of the fringe, of 1 or lower, as determined by optical surface profilometry diameter ratio, which is ratio of the diameter of the hollow center at half height and the diameter A film, sheet or coating according to claim 1, wherein the fringes have a hollow
- 100 or lower, as determined by optical surface profilometry. Index, which is calculated by multiplying the hollow depth ratio with the hollow diameter ratio, of A film, sheet or coating according to claim 1, wherein the fringes have a Hollowness
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- biaxially oriented polypropylene film. layers is a fringed layer and at least one of the inner layers is an oriented film, preferably a 13. The film, sheet or coating according to claim 10, wherein at least on of the outer
- foamed layer. 14. The film, sheet or coating according to claim 13, wherein at least one of the layers is a
- 15. The film, sheet, or coating according to claim 1, wherein at least one layer is elastic
- 16. The film, sheet, or coating according to claim 1, which is oriented.
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- 21. The composite according to claim 20 which is a laminate.
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- 27. The article of manufacture according to claim 22 which has a soft, textile-like touch.

- 28. The article of manufacture according to claim 22 which is water repellant.
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- 31. The article of manufacture according to claim 22 which is heat resistant.
- claim 1, said process comprising 32. A process for making the mono-layer or multi-layer film, sheet, or coating according to
- forming a precursor film, sheet, or coating with a surface characterized by a pattern of peaks and valleys in a continuous compression molding process, and
- subjecting said precursor to mechanical treatment comprising the application of a matrix surface under conditions allowing the formation of a fringed surface tractive force which is applied during release of the film, sheet or coating from a microstructure.
- 170 degrees relative to the matrix surface. the Vicat softening point of the thermoplastic material and at a release angle of between 20 and peeling the film, sheet, or coating off the matrix surface at a temperature which is at or below 33. The process according to claim 32, wherein application of the tractive force comprises
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- 38. Use of the film, sheet or coating according to claim 1 for packaging.
- carrying, capturing or storing properties. 39. Use of the film, sheet or coating according to claim 1 in an article with enhanced
- anti-skid properties. 40. Use of the film, sheet or coating according to claim 1 to make an article which has
- enhanced heat resistance. 41. Use of the film, sheet or coating according to claim 1 to make an article which has
- matte surface appearance. 42. Use of the film, sheet or coating according to claim 1 to make an article which has a
- 43. Use of the film, sheet or coating according to claim 1 as a filtration medium.
- 44. Use of a thermoplastic material to make the film, sheet or coating according to claim

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- biodegradable[, preferably poly(lactide)]. (Amended) Use of a thermoplastic material according to claim [46] 44 which is
- 46. (New) The article of manufacture according to claim 22, which is an ostomy bag.
- material is a poly(lactide). 47. (New) Use of a thermoplastic material according to claim 44 which thermoplastic

FRINGED SURFACE STRUCTURES OBTAINABLE IN A COMPRESSION MOLDING PROCESS

FIELD OF THE INVENTION

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and a device to make such film, sheet, or coating, uses thereof, as well as articles of manufacture made therefrom. has or comprises a fringed surface microstructure. multi-layer film, sheet, or coating characterized in that it comprises at least one layer which This invention relates to a surface having a fringed microstructure and to a mono-layer or The invention also provides a process

BACKGROUND OF THE INVENTION

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surface-structured films include apertured films and non-apertured films or garments. To improve aesthetics, for example of a plastic film, various techniques of touch, and good optics are desirable for numerous applications, including hygienic products surface modification resulting in various surface structures have been employed. Known Plastic articles and items with a pleasant haptic appearance, such as a textile-like

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through the surface feature, e.g. the protruberance. of the film, irrespective of any surface structure or pattern, and films which are apertured Surface-structured apertured films include films wherein the aperture is in the plane

exhibiting a pattern of discrete volcano-like aberrations, the end of which includes at least film is supported on a fine mesh woven wire support member. The high pressure fluid jet a jet of high pressure liquid on the exposed surface of a web of flat polymeric film while said one micro-aperture at its point of maximum amplitude. The films are produced by impinging between the intersecting woven wire filaments comprising the support member and which causes micro-aperturing of those portions of the web which coincide with the interstices correspond to the surface aberrations after removal of the web. For example, US Patent No. 4,629,643 provides a micro-apertured polymeric web

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apertured film. comprises a woven or non-woven fibrous material and an elastomeric three-dimensional US Patent No. 5,733,628 discloses a film laminate wherein the carrier material

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Non-apertured surface structured films include films with solid or with hollow surface

discloses extruded films which due to melt fracture show a rough surface texture by taking advantage of the usually undesired phenomenon of melt fracture. The patent According to U.S. Patent No. 5,814,413 surface-texturing of a polymer film is accomplished

projections is peeled off the surface. solidification of the thermoplastic material, the semi-finished product covered with fiber-like air from the cavities and enabling these to be filled with the thermoplastic material. After millimeters. The roller surface is exposed to a vacuum from the outside thus removing the process, the thermoplastic material is extruded in the molten state onto a moving belt or surface-structured, sheet-like semi-finished product made from thermoplastic polymers. The roller surface which is covered with cavities or borings having a depth of between 2 and 4 fibrous projections which can be longer than 3 millimeters. According to the disclosed resulting product is reported to have a velour-like or velvety surface with numerous solid International Patent Application WO 97/02128 provides a process for producing

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brushing, knife-coating and/or shear pinching pile consisting of solid protuberances and elongating the protruberances by combing surface-structured, film-like semi-finished product from a thermoplastic comprising forming a International Patent Application WO 99/47339 describes a method for producing

two major surfaces of an oriented film, embossing the softened surface(s), and cooling the resulting embossed oriented thermoplastic film. In order to maintain orientation the combined steps of softening, embossing and cooling should occur within a second embossed oriented film. Said method discloses the steps of softening at least one of the International Patent Application WO 99/16608 discloses a method for making an

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microfibers, tapered microfibers, microfibers having an expanded cross-sectional shape and area and to impart a cloth-like feel. microfibers having a high aspect ratio. The microfibers are reported to increase the surface having a plurality of solid microfibers which may have a variety of forms, such as frayed-end International Patent Application WO 99/06623 provides a unitary polymer substrate

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naps which may be stretched, e.g. by brushing. as a structured film. The disclosed structure is a pile comprised of solid projections and surface-structured, film-like semifinished product made of a thermoplastic which is applied a surface covered with fine cavities. The solidified plastic is removed from the surface International Patent Application WO 00/16965 relates to a method for producing a

96/13979 and US Patent No. 5,192,484. disclosed, for example, in US Patent No. 4,463,045, International Patent Application WO Films with bubble-like surface features which are hollow from the bottom are

a suction cup geometric configuration. U.S. Patent No. 5,792,411 suggests replicated articles with surface structures which

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release or carrier properties, controlled thermal and barrier performance, as well as any surface area, variability of the surface appearance, anti-slip behavior, controlled storage, representative properties desirable for plastic articles include, for example, increased excellent aesthetics, as reflected in a textile-like haptic appearance and low gloss. Further combination thereof, There still is the need for plastic articles showing improved properties, particularly

advantageous performance attributes depending on the intended end-use application(s). desired combination of the above-mentioned properties, and may afford additional particular object of the present invention to provide a plastic article, which displays any providing a plastic article having a frictional behavior indicating anti-slip properties. It is a article with an imprintable surface. The present invention also addresses the problem of combination with a mat appearance. It is another object to provide a plastic article having an present invention to provide a plastic article having a soft, velvety and cloth-like touch in attribute(s) and can be produced in a cost effective way. In particular, it is one object of the surface structures which can be specifically designed to meet the desired performance present invention to provide surface-structured plastic films, sheets or coatings with hollow increased surface area. It is yet a further object of the present invention to provide a plastic It is an object of the present invention to meet these needs. It is an object of the

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the intended end-use application(s). coating, which is characterized by a distinct morphology. Such morphology is reflected in invention can be tailored to satisfy the particular requirements of the production process and further layers affording certain additional functions, for example, heat sealability, bulk or the presence of at least one layer having a fringed surface microstructure and, optionally, of mechanical properties. The objects of the present invention are achieved by providing a film, sheet, or The morphology and composition of the film, sheet or coating of the

SUMMARY OF THE INVENTION

may be discernible with the naked eye or on appropriate magnification. entirety, or in part, e.g. in one area or in several areas. The fringed surface microstructure centimeter, or higher. The layer may be covered by the fringed surface microstructure in its are at least partially hollow. The density of the fringes is at least 1000 per square thermoplastic polymeric material and which is characterized by a fringed surface The film, sheet or coating of the invention comprises at least one layer which is a The present invention pertains to a mono-layer or multi-layer film, sheet, or coating. The fringes of said microstructure are non-perforated crater-like peaks which

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least one layer has a fringed surface microstructure. multi-layer film, sheet, or coating based on a thermoplastic polymeric material wherein at Another aspect of the invention relates to a composite comprising a mono-layer or

polymeric material wherein at least one layer has a fringed surface microstructure made from a mono-layer or multi-layer film, sheet, or coating based on a thermoplastic Another aspect of the invention relates to an article of manufacture comprising or

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fringed surface microstructure, said process comprising: coating based on a thermoplastic polymeric material wherein at least one layer has a Another aspect of the invention relates to a process for making the film, sheet, or

providing a precursor film, sheet, or coating with a surface characterized by a pattern of peaks and valleys, and

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 treating said precursor such as to create a surface having a fringed microstructure Preferably, the treatment is mechanical in nature.

which benefit from the advantageous properties and performance attributes provided by the one layer has or comprises the fringed surface microstructure in applications or articles sheet, or coating which is based on a thermoplastic polymeric material and wherein at least fringed surface microstructure Yet another aspect of the invention relates to the use of a mono- or multilayer film,

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BRIEF DESCRIPTION OF THE DRAWINGS

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its three-dimensional structure FIG. 1 shows an exemplary individual fringe and selected parameters characterizing

having a fringed microstructure FIG. 2 shows an example of a device suitable to make a film, sheet or coating

coating having the fringed surface microstructure. coating is stripped from the matrix roll in a calander suitable to make the film, sheet or 3 illustrates a method to determine the release angle at which the film, sheet or

DETAILED DESCRIPTION OF THE INVENTION

Basic Definitions

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generally exclude the fringe height. ribbons. The article has a flat form and a thickness of about 10 mils (250 microns (μm) or micrometer), or less. Thickness data for a film having a fringed surface microstructure The term "film" as used herein refers to a thin article and includes strips, tapes, and

than about 10 mils (250 microns). The term "sheet" as used herein refers to a flat article having a thickness of greater

7 contour. For example, a film, profile or molding may be coated or comprise a coating. Also, such as, for example, but not limited to, polyethylene, polypropylene, paper, glass, ceramic, the substrate surface can be made from a thermoplastic or a non-thermoplastic material substrate material. The substrate material (or its surface) can be of any shape, form or cardboard, foil, wood or wood-based materials, such as particleboard or fiberboard, and metal, such as copper. The term "coating" as used herein refers to a material applied over or on top of a

20 five, six, seven or more layers. layer film") as used herein indicates a film, sheet, or coating consisting of two, three, four, The term "multi-layer film, sheet, or coating." (including each individually e.g. "multi-

layer of the structure is foamed and has a density less than the non-foamed polymer. film") as used herein refers to a mono-layer or multi-layer structure wherein at least one The term "foamed film, sheet, or coating" (including each individually e.g. "foamed

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surface structure, (including, for example, but not limited to, a fabric or a laminated structure article or material comprising at least one film, sheet, or coating layer having a fringed may comprise, for example foil or paper). The term "composite" as used herein refers to a multi-layer or multi-component

obtainable by polymerizing one or more monomers. The generic term "polymeric compound" The term "polymeric material" as used herein refers to a polymeric compound

prepared from only one monomer, and an interpolymer as defined hereinafter or "polymer" is intended to include a homopolymer, usually employed to refer to polymers

The term "comprising" as used herein means "including"

monomers, and polymers prepared from more than two different monomers, such as polymerization of at least two monomers. The generic term interpolymer thus embraces the terms copolymer, usually employed to refer to polymers prepared from two different The term "interpolymer" as used herein refers to polymers prepared by the

to an aliphatic or cyclo-aliphatic alpha-olefin having at least 3, preferably 3 to 20 carbon Unless specified otherwise, the term "alpha-olefin" ("lpha-olefin") as used herein refers

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stand for "cubic centimeters". equal to the numerical value which follows this expression. The expressions "cc" or "ccm" expression "up to" when used to specify a numerical range includes any value less than or Unless indicated to the contrary, all parts, percentages and ratios are by weight. The

Thermoplastic Materials

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thermoplastic polymeric materials, including, for example, latex. Preferred thermoplastic polymers, polyvinylidene chloride polymers and polyesters, including certain elastomers and block polymers. Semicrystalline thermoplastic materials and blends thereof are preferred polycarbonates, polyamides, polyethers, thermoplastic phenoxy resins, polyvinyl chloride of polyolefins, poly(lactide), alkenyl aromatic polymers, thermoplastic polyurethanes polymeric materials are semicrystalline polymers, amorphous polymers, or blends thereof. Advantageously, suitable thermoplastic polymers may be selected from the group consisting The film, sheet, or coating of the invention is based on or made from one or more

polypropylene, polybutene and polyisoprene, and their interpolymers ethylene homopolymer and interpolymer, aliphatic alpha-olefin homopolymers, such as Suitable polyolefins include, for example, ethylene-based polymers, including

thermoplastic polymers, each having many members. They are prepared by density polyethylene (HDPE), and ethylene interpolymers are known classes of Ethylene homopolymers, for example low density polyethylene (LDPE) and high

or more sites of ethylenic unsaturation, etc., using known copolymerization reactions and one or more vinyl- or diene-based comonomers, for example, α -olefins of 3 to about 20 carbon atoms, vinyl esters, vinyl acids, styrene-based monomers, monomers containing two homopolymerizing ethylene or interpolymerizing (for example, copolymerizing) ethylene with

homogeneously branched (homogeneous) polymers and heterogeneously branched (heterogeneous) polymers Ethylene (based) polymers suitable for use in the present invention include both

such as is the case when an ethylene polymer having a crystallinity of less than about 36 differential scanning calorimetry (DSC). The single melting peak may be relatively broad, essentially single melting (point) peak between -30°C and 150°C, as determined by interpolymer. Homogeneous ethylene polymers generally are characterized as having an all of the interpolymer molecules have the same ethylene/comonomer ratio within that comonomer is randomly distributed within a given interpolymer molecule and substantially ethylene polymer having a crystallinity of at least about 36 percent is employed percent is employed. "Homogeneous" polymers encompass ethylene-based interpolymers in which any The single melting peak may be sharp, such as is the case when an

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about 3.5 (when the density of the interpolymer is less than about 0.960 g/cc), more weight average molecular weight to number average molecular weight ($M_W/M_{
m p}$), is less than ethylene polymers. Preferably, the molecular weight distribution defined as the ratio of molecular weight distribution (MWD) as compared to corresponding heterogeneous preferably less than about 3.0. Typically, homogeneous ethylene polymers will also have a relatively narrow

such SCBDI (Short Chain Branch Distribution Index) or (CDBI Composition Distribution techniques known in the art, such as, for example, temperature rising elution fractionation Branch Index). The SCBDI of a polymer is readily calculated from data obtained from reflected in a narrow composition distribution, which can be expressed using parameters a comonomer content within 50 percent of the median total molar comonomer content. The herein by reference. CDBI is defined as the weight percent of the polymer molecules having in U.S. Patent 5,089,321 (Chum et al.), the disclosures of all of which are incorporated Science, Poly. Phys. Ed., Vol. 20, p. 441 (1982), in U.S. Patent 4,798,081 (Hazlitt et al.), or (abbreviated herein as "TREF") as described, for example, in Wild et al, Journal of Polymer In addition or in the alternative, the homogeneity of the ethylene-based polymers

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polymers used in the present invention is typically greater than about 50 percent SCBDI or CDBI for the homogeneous linear and substantially linear ethylene/alpha-olefin

linear homogeneous ethylene polymers. into two broad categories, the linear homogeneous ethylene polymers and the substantially The homogeneous ethylene polymers that can be used in the present invention fall Both are known.

resins and by Exxon Chemical Company as EXACT™ and EXCEED™ resins. polymers include, for example, those sold by Mitsui Petrochemical Industries as TAFMER™ linear ethylene polymers. Commercially available examples of homogeneous linear ethylene catalysts, such as catalyst systems based on hafnium, for the preparation of homogeneous Ewen et al. and U.S. Patent No. 5,218,071 to Tsutsui et al. disclose the use of metallocene can be prepared in conventional polymerization processes using Ziegler-type catalysts such exemplified in U.S. Patent No. 3,645,992 to Elston, homogeneous linear ethylene polymers for example, zirconium and vanadium catalyst systems. U.S. Patent No. 4,937,299 to Homogeneous linear ethylene polymers have long been commercially available.

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having long chain branching. The substantially linear ethylene polymers (SLEPs) are homogeneous polymers

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branches/1000 total carbons to about 1 long chain branches/1000 total carbons about 1 long chain branched/1000 total carbons, and especially from about 0.3 long chain total carbons, more preferably from about 0.05 long chain branches/1000 total carbons about 0.01 long chain branches/1000 total carbons to about 1 long chain branches/1000 includes both backbone and branch carbon atoms). Preferred polymers are substituted with carbons to about 3 long chain branches/1000 total carbons (wherein "total carbons" ethylene polymer is substituted, on average, with about 0.01 long chain branches/1000 total The term "substantially linear ethylene polymer" as used herein means that the bulk

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to about 3 long chain branches/1000 total carbons branching in the bulk polymer is at least an average of from about 0.01/1000 total carbons have at least enough molecules with long chain branching such that the average long chain "polymer" or "bulk polymer" refers, in the conventional sense, to the polymer as formed in a reactor. For the polymer to be a "substantially linear ethylene polymer", the polymer must As used herein, the term "backbone" refers to a discrete molecule, and the term

polymerization process as a mixture of polymer molecules and, for substantially linear The term "bulk polymer" as used herein means the polymer which results from the 30

the melt fracture properties) as described herein below and elsewhere in the literature. long chain branching content of the bulk polymer positively affects the melt rheology (i.e., formed during polymerization. It is understood that, for the substantially linear polymers, not all molecules have long chain branching, but a sufficient amount do such that the average as molecules having long chain branching. Thus a "bulk polymer" includes all molecules ethylene polymers, includes molecules having an absence of long chain branching as well

six (6) carbons in length. branches of at least seven (7) carbons in length, but it also has short chain branches of only example, a substantially linear ethylene/1-octene polymer has backbones with long chain residue of the comonomer after it is incorporated into the polymer molecule backbone. For branching (SCB) is defined herein as a chain length of the same number of carbons in the carbon (atom) less than the number of carbons in the comonomer, whereas short chain Long chain branching (LCB) is defined herein as a chain length of at least one (1)

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and as such, this analytical technique cannot distinguish between a seven (7) carbon the same length as the length of the polymer backbone. branch and a seventy (70) carbon branch. The long chain branch can be as long as about cannot determine the length of a long chain branch in excess of about six (6) carbon atoms However as a practical matter, current ¹³C nuclear magnetic resonance spectroscopy homopolymers, it can be quantified using the method of Randall, (Rev. Macromol.Chem. nuclear magnetic resonance (NMR) spectroscopy and to a limited extent, e.g. for ethylene Phys., C29 (2&3), p. 285-297), the disclosure of which is incorporated herein by reference. Long chain branching can be distinguished from short chain branching by using 13C

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characteristics are determined by gel permeation chromatography and intrinsic viscosity average number of long chain branches per molecule and $M_{f W}$ is the weight average frequency (LCB) can be represented by the equation LCB = $\mathrm{b/M_W}$ wherein b is the weight known techniques useful for quantifying or determining the presence of long chain branches determine the length of a long chain branch in excess of six carbon atoms, there are other methods, respectively. molecular weight. No. 4,500,648, incorporated herein by reference, teaches that long chain branching in ethylene polymers, including ethylene/1-octene interpolymers. Although conventional 13C nuclear magnetic resonance spectroscopy cannot The molecular weight averages and the long chain branching For example, US Patent

theories have been well documented in the literature. See, e.g., Zimm, G.H. and disclosures of both of which are incorporated by reference. Polymer Characterization, John Wiley & Sons, New York (1991) pp. 103-112, the Stockmayer, W.H., J. Chem. Phys., 17, 1301 (1949) and Rudin, A., Modern Methods of (GPC-DV). The use of these techniques for long chain branch detection and the underlying permeation chromatography coupled with a low angle laser light scattering detector (GPCbranches in ethylene polymers, including ethylene/1-octene interpolymers are gel LALLS) and gel permeation chromatography coupled with a differential viscometer detector Two other useful methods for quantifying or determining the presence of long chain

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of long chain branches in substantially linear ethylene homopolymer samples measured substantially linear ethylene polymers. In particular, deGroot and Chum found that the level using the Zimm-Stockmayer equation correlated well with the level of long chain branches indeed a useful technique for quantifying the presence of long chain branches in measured using 13C NMR. Society (FACSS) in St. Louis, Missouri, presented data demonstrating that GPC-DV is October 4, 1994 conference of the Federation of Analytical Chemistry and Spectroscopy A. Willem deGroot and P. Steve Chum, both of The Dow Chemical Company, at the

account for the molecular weight increase attributable to octene short chain branches substantially linear ethylene/octene copolymers. showed that GPC-DV may be used to quantify the level of long chain branches in molecular weight increase attributable to 1-octene short chain branches, deGroot and Chum knowing the mole percent octene in the sample. By deconvoluting the contribution to hydrodynamic volume of the polyethylene samples in solution and, as such, one can Further, deGroot and Chum found that the presence of octene does not change the

polyethylene (LDPE) and are clearly distinct from ethylene polymers produced using long chain branching aspects (but not the extent of long branching) of substantially linear such as hafnium and vanadium complexes Ziegler-type catalysts such as titanium complexes and ordinary homogeneous catalysts ethylene polymers are comparable to that of high pressure, highly branched low density log(GPC Weight Average Molecular Weight) as determined by GPC-DV illustrates that the DeGroot and Chum also showed that a plot of log(l2, melt index) as a function of

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long chain branching is manifested as enhanced rheological properties which are quantified For substantially linear ethylene polymers, the empirical effect of the presence of

and expressed in terms of gas extrusion rheometry (GER) results and/or melt flow, I_{10}/I_2 , increases

polyolefin elastomers (POEs). polyolefin plastomers (POPs), and from DuPont Dow Elastomers, L.L.C. as ENGAGETM as polymers made by the INSITE™ Process and Catalyst Technology, such as AFFINITY™ incorporated herein by reference. SLEPs are available from The Dow Chemical Company 5,272,236, US Patent No. 5,278,272, and US Patent No. 5,665,800, each of which is invention are a unique class of compounds that are further defined in US Patent No. The substantially linear ethylene polymers suitable for the purpose of the present

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substantially linear ethylene polymers are characterized as essentially lacking a measurable also differ from the class of heterogeneously branched ethylene polymers in that more preferably greater than or equal to 90 percent. SCBDI greater than or equal to 50 percent, preferably greater than or equal to 70 percent, homogeneously branched polymers; that is, substantially linear ethylene polymers have a in US Patent No. 4,076,698, in that substantially linear ethylene interpolymers are polyethylene (HDPE) made, for example, using the technique disclosed by Anderson et al. low density polyethylene, linear low density polyethylene (LLDPE) or high density branched traditional Ziegler polymerized linear ethylene interpolymers (for example, ultra also differ significantly from the class of polymers known conventionally as heterogeneously homogeneously branched linear ethylene polymers. Substantially linear ethylene polymers linear polymer backbone in the conventional sense of the term "linear" as is the case for above. As an important distinction, substantially linear ethylene polymers do not have a conventionally known as homogeneously branched linear ethylene polymers described high density or crystalline polymer fraction as determined using a temperature rising elution fractionation technique Substantially linear ethylene polymers differ significantly from the class of polymers Substantially linear ethylene polymers

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characterized as having The substantially linear ethylene polymer for use in the present invention can be

(a) melt flow ratio, $1_{10}/1_2 \cdot 5.63$

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ਰ permeation chromatography and defined by the equation: a molecular weight distribution, M_W/M_D , as determined by gel

 $(M_W/M_n) \le (I_{10}/I_2) - 4.63$

(c) a gas extrusion rheology such that the critical shear rate at onset of

ethylene polymer are measured at the same melt temperature using a gas extrusion respective critical shear rates of the substantially linear ethylene polymer and the linear M_W/M_D within ten percent of the substantially linear ethylene polymer and wherein the comprise the same comonomer or comonomers, the linear ethylene polymer has an ${
m l}_2$ and polymer, wherein the substantially linear ethylene polymer and the linear ethylene polymer greater than the critical shear rate at the onset of surface melt fracture for a linear ethylene surface melt fracture for the substantially linear ethylene polymer is at least 50 percent

and 150°C, and (d) a single differential scanning calorimetry, DSC, melting peak between -30°C

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(e) a short chain branching distribution index greater than 50 percent

described by M. Shida, R.N. Shroff and L.V. Cancio in Polymer Engineering Science, Vol. by Van Nostrand Reinhold Co. (1982) on pp. 97-99, the disclosures of both of which are fracture as well as other rheology properties such as "rheological processing index" (PI), is 17, No. 11, p. 770 (1977) and in Rheometers for Molten Plastics by John Dealy, published performed using a gas extrusion rheometer (GER). The gas extrusion rheometer is incorporated herein by reference. Determination of the critical shear rate and critical shear stress in regards to melt

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units from the following equation: pressure of 2500 psig using a 0.0296 inch (752 micrometers) diameter (preferably a 0.0143 having an entrance angle of 180°. inch diameter die for high flow polymers, e.g. 50 - 100 $m l_2$ melt index or greater), 20:1 L/D die The processing index (PI) is measured at a temperature of 190°C, at nitrogen The GER processing index is calculated in millipoise

 $PI = 2.15 \times 10^6 \text{ dyne/cm}^2/(1000 \times \text{shear rate})$

rate is the shear rate at the wall as represented by the following equation: wherein: $2.15 imes 10^6$ dyne/cm 2 is the shear stress at 2500 psi, and the shear

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32 Q'/ (60 sec/min)(0.745)(Diameter X 2.54 cm/in)³, wherein:

Q' is the extrusion rate (gms/min),

0.745 is the melt density of polyethylene (gm/cm³), and Diameter is the orifice diameter of the capillary (inches)

The PI is the apparent viscosity of a material measured at apparent shear stress

 $2.15 \times 10^{6} \text{ dyne/cm}^{2}$

within ten percent of the substantially linear ethylene polymer. of that of a conventional linear ethylene polymer having an $m I_2$, $m M_w/M_n$ and density each For substantially linear ethylene polymers, the PI is less than or equal to 70 percent

irregularities may be broadly classified into two main types: surface melt fracture and gross or GER test apparatus previously described. According to Ramamurthy in Journal of fracture phenomena over a range of nitrogen pressures from 5250 to 500 psig using the die Rheology, 30(2), 337-357, 1986, above a certain critical flow rate, the observed extrudate An apparent shear stress vs. apparent shear rate plot is used to identify the melt

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٦ ٦ 10 polymers is at least 50 percent greater than the critical shear rate at the onset of surface Preferably, the critical shear stress at onset of surface melt fracture for the substantially at which the surface roughness of extrudate can only be detected by 40x magnification. linear ethylene polymers of the invention is greater than about 2.8 \times 10 6 dyne/cm 2 melt fracture of a linear ethylene polymer having about the same I $_2$ and M $_{
m W}$ /M $_{
m n}$. The critical shear rate at onset of surface melt fracture for the substantially linear ethylene the onset of surface melt fracture is characterized at the beginning of losing extrudate gloss detail from loss of specular gloss to the more severe form of "sharkskin". In this disclosure, Surface melt fracture occurs under apparently steady flow conditions and ranges in

linear ethylene polymers used in the invention, the critical shear stress at onset of gross onset of gross melt fracture (OGMF) will be used herein based on the changes of surface regular (alternating rough and smooth, helical, etc.) to random distortions. For commercial melt fracture is preferably greater than about 4 x 10⁶ dyne/cm² roughness and configurations of the extrudates extruded by a GER. For the substantially The critical shear rate at onset of surface melt fracture (OSMF) and critical shear stress at acceptability, (e.g., in blown film products), surface defects should be minimal, if not absent. Gross melt fracture occurs at unsteady flow conditions and ranges in detail from

such as phenols, hindered phenols, phosphites or phosphonites, preferably a combination index and melt fracture tests, substantially linear ethylene polymers do contain antioxidants of a phenol or hindered phenol and a phosphite or a phosphonite more than 20 ppm aluminum catalyst residue. Preferably, however, for the processing substantially linear ethylene polymers are tested without inorganic fillers and do not have For the processing index determination and for the GER melt fracture determination,

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percent by weight solutions of the samples are prepared for injection. The flow rate is about about 100 microliters. 1.0 milliliters/minute, unit operating temperature is about 140°C and the injection size is 10^3 , 10^4 , 10^5 and 10^6 Å. The solvent is 1,2,4-trichlorobenzene, from which about 0.3 columns are supplied by Polymer Laboratories and are commonly packed with pore sizes of unit equipped with a differential refractometer and three columns of mixed porosity. The permeation chromatography (GPC) on a Waters 150 °C high temperature chromatographic The molecular weight distributions of ethylene polymers are determined by gel

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herein by reference) to derive the following equation: Science, Polymer Letters, Vol. 6, p. 621, 1968, the disclosure of which is incorporated polyethylene and polystyrene (as described by Williams and Ward in Journal of Polymer molecular weights are determined by using appropriate Mark-Houwink coefficients for Laboratories) in conjunction with their elution volumes. The equivalent polyethylene deduced by using narrow molecular weight distribution polystyrene standards (from Polymer The molecular weight determination with respect to the polymer backbone is

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$$M_{\text{polyethylene}} = a * (M_{\text{polystyrene}})b.$$

calculated in the usual manner according to the following formula: In this equation, a = 0.4316 and b = 1.0. Weight average molecular weight, M_W , is

$$M_{j} = (\sum w_{i}(M_{i}^{i}))^{i};$$

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wherein w, is the weight fraction of the molecules with molecular weight M, eluting from the GPC column in fraction i and j=1 when calculating M_{w} and j=-1 when calculating M_{p} .

branched linear ethylene polymers, the melt flow ratio (110/12) of substantially linear typically less than about 3.5). Surprisingly, unlike homogeneously and heterogeneously ethylene polymers can be varied essentially independently of the molecular weight despite having a relatively narrow molecular weight distribution (that is, the M_W/M_D ratio is distribution, Mw/Mn. Substantially linear ethylene polymers are known to have excellent processability,

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ethylene polymers include constrained geometry catalysts as disclosed in U.S. application number 07/545,403, filed July 3, 1990; U.S. application number 07/758,654, filed September 12, 1991; U.S. Patent No. 5,132,380; U.S. Patent No. 5,064,802; U.S. Patent Suitable constrained geometry catalysts for manufacturing substantially linear

of which are incorporated herein by reference 5,532,394; U.S. Patent No. 5,494,874; and U.S. Patent No. 5,189,192, the teachings of all No. 5,470,993; U.S. Patent No. 5,453,410; U.S. Patent No. 5,374,696; U.S. Patent No

Patent No. 5,272,236; US Patent No. 5,278,272 and US Patent No. 5,665,800, especially catalysts taught in U.S. Patent No. 5,026,798, which is incorporated herein by reference, methods are also described in PCT/US 92/08812 (filed October 15, 1992). with strict attention to the requirement of continuous polymerization. Such polymerization so long as the polymerization conditions substantially conform to those described in US are also believed to be suitable for use in preparing the polymers of the present invention, reference. Further, the monocyclopentadienyl transition metal olefin polymerization Suitable catalyst complexes may also be prepared according to the teachings and the patents issuing therefrom, all of which are incorporated herein by

delocalized, substituted pi-bonded moiety, only one thereof for each metal atom of the substituent, and provided further that for such complexes comprising more than one the Elements and a delocalize β -bonded moiety substituted with a constrain-inducing complex is a cyclic, delocalized, substituted pi-bonded moiety. The catalyst further complex containing a similar pi-bonded moiety lacking in such constrain-inducing and the center of at least one remaining substituent is less than such angle in a similar angle at the metal between the centroid of the delocalized, substituted pi-bonded moiety moiety, said complex having a constrained geometry about the metal atom such that the complex comprising a metal of groups 3-10 or the Lanthanide series of the Periodic Table of comprises an activating cocatalyst. The foregoing catalysts may be further described as comprising a metal coordination

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US Patent No. 4,960,878; and US Patent No. 4,544,762, the disclosures of all of which are 5,086,024; US Patent No. 5,041,585; US Patent No. 5,041,583; US Patent No. 5,015,749; Patent No. 5,041,584, the disclosure of which is incorporated herein by reference. compounds. So called modified methyl aluminoxane (MMAO) is also suitable for use as especially methyl aluminoxane, as well as inert, compatible, noncoordinating, ion forming Aluminoxanes can also be made as disclosed in US Patent No. 5,218,071; US Patent No. incorporated herein by reference Suitable cocatalysts for use herein include polymeric or oligomeric aluminoxanes, One technique for preparing such modified aluminoxane is disclosed in US

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aluminum. In order to measure the bulk polymer properties (e.g. PI or melt fracture), polymer is preferably in the range of from about 0 to about 20 ppm aluminum, especially 0520732, the disclosure of which is incorporated herein by reference however, are inert, noncoordinating, boron compounds such as those described in EP-Aaqueous HCl is used to extract the aluminoxane from the polymer. Preferred cocatalysts, from about 0 to about 10 ppm aluminum, and more preferably from about 0 to about 5 ppm polymerization, are preferably used such that the catalyst residue remaining in the (finished) Aluminoxanes, including modified methyl aluminoxanes, when used in the

can be operated in series or in parallel, with at least one constrained geometry catalyst sufficient to produce the interpolymers having the desired properties. which is incorporated herein by reference) at a polymerization temperature and pressure multiple reactor configuration as described in US Patent No. 3,914,342, the disclosure of herein by reference), but can also be produced using multiple reactors (e.g., using a 93/07187, WO 93/07188, and WO 93/07189, the disclosure of each of which is incorporated controlled polymerization process using at least one reactor (e.g., as disclosed in WO employed in at least one of the reactors. Substantially linear ethylene are produced via a continuous (as opposed to a batch) The multiple reactors

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the like. Preferably, a continuous loop-reactor solution polymerization process is used to known in the art including, but not limited to, a tank reactor(s), a sphere reactor(s), a such as the method disclosed in EP-A-416,815, the disclosure of which is incorporated slurry, or gas phase polymerization in the presence of a constrained geometry catalyst, operated partially or completely adiabatically, nonadiabatically or a combination of both and recycling loop reactor(s) or combinations thereof and the like, any reactor or all reactors herein by reference. The polymerization can generally be performed in any reactor system manufacture the substantially linear ethylene polymer used in the present invention Substantially linear ethylene polymers can be prepared via the continuous solution,

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solution, slurry, gas phase or other process conditions may be employed if desired to 250°C and pressures from atmospheric to 1000 atmospheres (100 MPa). Suspension, linear ethylene polymers may be accomplished at conditions well known in the prior art for Ziegler-Natta or Kaminsky-Sinn type polymerization reactions, that is, temperatures from 0 In general, the continuous polymerization required to manufacture substantially

used in a homogeneous (i.e., soluble) manner. It will, of course, be appreciated that the support may be employed in the polymerization, but preferably the catalysts are

active catalyst system forms in situ if the catalyst and the cocatalyst components thereof are form the active catalyst in a separate step in a suitable solvent prior to adding the same to added directly to the polymerization process and a suitable solvent or diluent, including the polymerization mixture. condensed monomer, is used in said polymerization process. It is, however, preferred to

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accordance with the definitions and preferences given hereinbelow. Copolymers of ethylene are interpolymers of ethylene with at least one C3-C20 lpha-olefin and/or C4-C18 diolefin, in and an α -olefin of C3-C20 carbon atoms are preferred. Preferably, the substantially linear ethylene polymers used in the present invention

H 0 15 ethylene interpolymers have a CDBI of about 50 % or less, indicating that such the density of the interpolymer is less than about 0.960 g/cc). Typically, heterogeneous homogeneous interpolymers, as reflected in a M_w/M_n ratio of greater than about 3.5 (when heterogeneous interpolymers will typically have a broader molecular weight distribution than distinct melting point peak greater than 115°C attributable to a high density fraction. Such interpolymers characterized as having a linear backbone and a DSC melting curve having a interpolymers are a mixture of molecules having differing comonomer contents and differing amounts of short chain branching. Heterogeneous ethylene-based polymers encompass ethylene/lpha-olefin

20 25 30 invention fall into two broad categories, those prepared with a free radical initiator at high units pendant from the polymer backbone. LDPE polymers generally have a density between about 0.910 and 0.935 g/cc. Ethylene polymers and copolymers prepared by the polyethylenes (LDPE) and are characterized by branched chains of polymerized monomer temperature and relatively low-pressure. The former are generally known as low density temperature and high pressure, and those prepared with a coordination catalyst at high and it contains relatively few branch chains relative to the various linear copolymers or having a density of about 0.941 to about 0.965 g/cc, is typically a homopolymer of ethylene, monomer units pendant from the backbone. High density polyethylene (HDPE), generally linear polymers because of the substantial absence of branch chains of polymerized use of a coordination catalyst, such as a Ziegler or Phillips catalyst, are generally known as ethylene and an α -olefin. HDPE is well known, commercially available in various grades, and may be used in this invention. The heterogeneous ethylene polymers that can be used in the practice of this

olefin content to reduce the density of the copolymer to that of LDPE. When the copolymer amount of ethylene, the greater the length of the lpha-olefin and the greater the amount of lphathe length of the α -olefin and the amount of such monomer in the copolymer relative to the well known in the art, the density of a linear ethylene/ α -olefin copolymer is a function of both 0.87 g/cc to about 0.91 g/cc. polyethylene (VLDPE). copolymers are known as ultra low density polyethylene (ULDPE) or very low density contains even more lpha-olefin, the density will drop below about 0.91 g/cc and these preferably 4 to 8 carbon atoms (for example, 1-butene, 1-octene, etc.), that has sufficient lpha-(LLDPE) is typically a copolymer of ethylene and an α -olefin of 3 to 12 carbon atoms, olefin present, the lower the density of the copolymer. Linear low density polyethylene preferably of 4 to 8 carbon atoms, are also well known and commercially available. As is Linear copolymers of ethylene and at least one lpha-olefin of 3 to 12 carbon atoms The densities of these linear polymers generally range from about

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al., which is incorporated herein by reference. catalyst, by processes such as are disclosed in U.S. Patent No. 4,076,698 to Anderson et ethylene and one or more optional α -olefin comonomers in the presence of a Ziegler Natta ethylene polymers can be prepared via the solution, slurry or gas phase polymerization of DOWLEX™ LLDPE polymers and as ATTANE™ ULDPE resins. Heterogeneous linear heterogeneous linear ethylene polymers are available from The Dow Chemical Company as catalysts are well known in the art, as are their methods of preparation. For example Both the materials made by the free radical catalysts and by the coordination

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tetrafluoroethylene, vinyl benzocyclobutane, 1,4-hexadiene, 1,7-octadiene, and combinations thereof include 1-propylene, 1-butene, 1-isobutylene, 1-pentene, 1-hexene, 4use as comonomers in a solution, gas phase or slurry polymerization process or methyl-1-pentene, 1-hexene, 1-heptene, 1-octene, NB or ENB, or mixtures thereof. More ethylidene norbornene (ENB)). Preferably, the α -olefin will be 1-butene, 1-pentene, 4methyl-1-pentene, 1-heptene and 1-octene, as well as other monomer types such as invention can be interpolymers of ethylene and at least one lpha-olefin. Suitable lpha-olefins for preferably, the lpha-olefin will be 1-pentene, 1-hexene, 1-heptene, 1-octene, or mixtures cycloalkenes, for example cyclopentene, cyclohexene, cyclooctene, norbornene (NB), and As indicated above, the ethylene polymers suitable for the purpose of the present Most preferably, the α -olefin will be 1-octene

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the terpolymer include: of such terpolymers are typically non-conjugated dienes having from 6 to 15 carbon atoms making ethylene lpha-olefin copolymers. The dienes suitable as monomers for the preparation Representative examples of suitable non-conjugated dienes that may be used to prepare this invention. Suitable lpha-olefins include the lpha-olefins described above as suitable for Ethylene/lpha-olefin/diene terpolymers may also be used as elastomeric polymers

- Ð Straight chain acyclic dienes such as 1,4-hexadiene, 1,5-heptadiene, and 1,6-octadiene;
- ᠑ branched chain acyclic dienes such as 5-methyl-1, 4-hexadiene, 3,7dimethyl-1-6-octadiene, and 3,7-dimethyl-1,7-octadiene

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- 0 single ring alicyclic dienes such as 4-vinylcyclohexene, 1-allyl-4isopropenyl-4-butenylcyclohexane; isopropylidene cyclohexane, 3-allylcyclopentene, 4-allylcyclohexene, and 1-
- 9 multi-ring alicyclic fused and bridged ring dienes such as dicyclopentadiene; alkenyl, alkylidene, cycloalkenyl and cycloalkylidene norbornenes such as 5norbornene, 5-ethylidene-2-norbornene, 5-cyclohexylidene-2-norbornene, 6, 6-dimethyl-2-norbornene, 5-propenyl-2-norbornene, 5-(3-cyclopentenyl)-2methylene-2-norbornene, 5-methylene-6-methyl-2-norbornene, 5-methylene-
- 20 octadiene, piperylene, 4-vinylcyclohexene, etc. dicyclopentadiene, 5-ethylidene-2-norbornene, 5-methylene-2-norbornene, 7-methyl-1, 6-The preferred dienes are selected from the group consisting of 1,4-hexadiene

make elastomeric polymer blends vulcanizable with the well known sulfur compound commercially available. Ethylene/ α -olefin/diene terpolymers are useful when it is desired ethylene, propylene and a non-conjugated diene (EPDM). Such terpolymers are vulcanization process The preferred terpolymers for the practice of the invention are terpolymers

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selected from the group consisting of vinyl esters of a saturated carboxylic acid wherein the acid moiety has up to 4 carbon atoms, unsaturated mono- or dicarboxylic acids of 3 to in this invention. alcohol having 1 to 8 carbon atoms, and mixtures thereof. Terpolymers of ethylene and carbon atoms, a salt of the unsaturated acid, esters of the unsaturated acid derived from an Ethylene/unsaturated carboxylic acid, salt and ester interpolymers may also be used These are interpolymers of ethylene with at least one comonomer

WO 01/05574

PCT/US00/19320

monoxide containing up to about 15 percent by weight of carbon monoxide can also be terpolymers of ethylene/vinyl acetate/carbon monoxide or ethylene/methyl acrylate/carbon detail in U.S. Patent 3,264,272, already incorporated herein by reference. In addition, neutralized copolymers of ethylene and the acids described above, are discussed in more these comonomers are also suitable. Ionomers, which are completely or partially

and its ionomers; ethylene/methacrylic acid and its ionomers; ethylene/methyl acrylate; ethylene/vinyl acetate (EVA) including, but not limited to, the stabilized EVA described in U. acrylate/methacrylic acid and its ionomers; ethylene/isobutyl acrylate/acrylic acid and its ethylene/isobutyl acrylate/methacrylic acid and its ionomers; ethylene/normal butyl S. Patent 5,096,955, which is incorporated herein by reference; ethylene/acrylic acid (EAA) EAA; ethylene/methyl acrylate; ethylene/isobutyl acrylate; and ethylene/methyl methacrylate ethylene/methyl acrylate/monoethyl maleate. Particularly suitable copolymers are EVA; ethylene/methacrylate/carbon monoxide; ethylene/normal butyl acrylate/carbon monoxide; acetate/acrylic acid and its ionomers; ethylene/vinyl acetate/carbon monoxide methacrylate; ethylene/vinyl acetate/methacrylic acid and its ionomers; ethylene/vinyl ionomers; ethylene/normal butyl acrylate/acrylic acid and its ionomers; ethylene/methyl ethylene/ethyl acrylate; ethylene/isobutyl acrylate; ethylene/normal butyl acrylate; these ethylene interpolymers are well known in the art and many are commercially Patent 4,379,190, which is incorporated herein by reference. The procedures for making be improved by certain combinations of these ethylene interpolymers described in U. S copolyers and mixtures thereof. Certain properties, such as tensile elongation, are taught to ethylene/isobutyl acrylate/carbon monoxide; ethylene/vinyl acetate/monoethyl maleate and Suitable ethylene/unsaturated carboxylic acid, salt and ester interpolymers include

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according to this invention. Such propylene based polymers are, for example ethylene and/or a C_4 - C_{20} alpha-olefin, including impact copolymers and polypropylene homopolypropylene and propylene interpolymers, such as copolymers of propylene with random copolymers Propylene based polymers are also suitable to make a film, sheet, or coating

polyolefin interpolymers comprising Further thermoplastic interpolymers suitable to practice the present invention are

- ت polymer units derived from at least one of ethylene and/or an alpha-olefin monomer; and
- € or vinylidene monomer, and vinyl or vinylidene monomers, or a combination of at least one aromatic vinyl polymer units derived from one or more vinyl or vinylidene aromatic monomers and/or one or more sterically hindered aliphatic or cycloaliphatic

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- ij optionally polymer units derived from one or more ethylenically unsaturated polymerizable monomer(s) other than those derived from i) and ii).
- 10 more of propylene, butene-1, 4-methyl-1-pentene, hexene-1 or octene-1. butene-1, 4-methyl-1-pentene, hexene-1 or octene-1 or ethylene in combination with one or olefins do not contain an aromatic moiety. Particularly suitable are ethylene, propylene, preferably from 3 to about 12, more preferably from 3 to about 8 carbon atoms. These lpha-Suitable α -olefins include, for example, α -olefins containing from 3 to about 20,
- exemplary interpolymer being ethylene/styrene/norbornene. such as norbornene and $\mathsf{C_1\text{-}C_{10}}$ alkyl or $\mathsf{C_6\text{-}C_{10}}$ aryl substituted norbornenes, with an Polymerizable ethylenically unsaturated monomer(s) include strained ring olefins

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represented by the following formula: Suitable vinyl or vinylidene aromatic monomers include, for example, those

$$\begin{array}{c}
Ar \\
(CH_2)_n \\
+ C = C(R^2)_2
\end{array}$$

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group or a phenyl group substituted with from 1 to 5 substituents selected from the group containing from 1 to about 4 carbon atoms, preferably hydrogen or methyl; Ar is a phenyl containing from 1 to about 4 carbon atoms, preferably hydrogen or methyl; each R² is wherein R' is selected from the group of radicals consisting of hydrogen and alkyl radicals styrene and lower alkyl- or halogen-substituted derivatives thereof. Preferred monomers include styrene, vinyl toluene, α -methylstyrene, t-butyl styrene, chlorostyrene, including all preferably from zero to 2, most preferably zero. Exemplary vinyl aromatic monomers consisting of halo, C_1 - C_2 -alkyl, and C_1 - C_2 -haloalkyl; and n has a value from zero to about 4, independently selected from the group of radicals consisting of hydrogen and alkyl radicals isomers of these compounds, and the like. Particularly suitable such monomers include

preferred aromatic vinyl monomer is styrene. halogenated styrenes, para-vinyl toluene or mixtures thereof, and the like. The most derivatives of styrene, such as for example, ortho-, meta-, and para-methylstyrene, the ring include styrene, lpha-methyl styrene, the lower alkyl-(C,-C $_{_4}$) or phenyl-ring substituted

to the formula: compounds", it is meant addition polymerizable vinyl or vinylidene monomers corresponding By the term "sterically hindered aliphatic or cycloaliphatic vinyl or vinylidene

$$\mathbb{R}^{1} \longrightarrow \mathbb{C} = \mathbb{C}(\mathbb{R}^{2})_{2}$$

7 10 wherein A' is a sterically bulky, aliphatic or cycloaliphatic substituent of up to 20 carbons, R' together form a ring system. from 1 to about 4 carbon atoms, preferably hydrogen or methyl; or alternatively R' and A' selected from the group of radicals consisting of hydrogen and alkyl radicals containing from 1 to about 4 carbon atoms, preferably hydrogen or methyl; each \mathbf{R}^2 is independently is selected from the group of radicals consisting of hydrogen and alkyl radicals containing

catalysts at a rate comparable with ethylene polymerizations. normally incapable of addition polymerization by standard Ziegler-Natta polymerization By the term "sterically bulky" it is meant that the monomer bearing this substituent is

butene-1, hexene-1 and octene-1 are not considered to be sterically hindered aliphatic Ethylene and alpha-olefins having a linear aliphatic structure such as propylene

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olefins including for example, α -olefins containing from 3 to about 20 carbon atoms such as derivatives of cyclohexene and substituted cyclohexenes, and 5-ethylidene-2-norbornene. cycloaliphatic vinyl or vinylidene compounds are the various isomeric vinyl-ring substituted substituted derivatives thereof, tert-butyl, norbornyl, and the like. Most preferred aliphatic or aliphatic groups such as cyclohexyl, cyclohexenyl, cyclooctenyl, or ring alkyl or aryl compounds are monomers in which one of the carbon atoms bearing ethylenic unsaturation sterically hindered aliphatic or cycloaliphatic vinyl or vinylidene compounds propylene, butene-1, 4-methyl-1-pentene, hexene-1 or octene-1 are not examples Especially suitable are 1-, 3-, and 4-vinylcyclohexene. Simple linear non-branched α is tertiarily or quaternarily substituted. Examples of such substituents include cyclic Preferred sterically hindered aliphatic or cycloaliphatic vinyl or vinylidene

preferred substantially random interpolymers are the so-called pseudo-random optionally variably sequenced. The preferred sequence is substantially random. The combination thereof. That is, the interpolymer product can be variably incorporated and substantially random, pseudo-random, random, alternating, diadic, triadic, tetradic or any No. 5,703,187 by Francis J. Timmers, both of which are incorporated herein by reference in interpolymers their entirety. The interpolymers comprising polymer units defined above under i), ii) and iii) can as described in EP-A-0 416 815 by James C. Stevens et al. and US Patent

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10 preferred are substantially random ethylene/styrene interpolymers. The preferred polyolefin interpolymer are ethylene/styrene interpolymers. Particularly

weight percent styrene and the production split between the two catalyst systems is 47/53 incorporates 22 weight percent styrene and the other catalyst system incorporates 48 styrene content of 36 weight percent is variably incorporated where one catalyst system incorporation or reactivity rates. For example, the interpolymer product having a total systems wherein during interpolymerization the catalyst systems are operated at different particularly an ethylene/styrene interpolymer, manufactured using at least two catalyst weight percentages The term "variably incorporated" as used herein refers to an interpolymer.

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interpolymers described in U.S. Patent 5,703,187, the disclosure of which is incorporated herein in its entirety by reference Representative "pseudo-random" interpolymers are the ethylene/styrene

where either the ethylene or the aliphatic alpha-olefin monomer or the sterically hindered vinylidene monomer or both can be repeated adjacent to one another. the chain such that there exist various combinations of ordering including blockiness, e.g. "Random" interpolymers are those in which the monomer units are incorporated into

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stereospecific structures (such as isotactic or syndiotactic) or in combinations of the general formula (AB), monomer occur in repeat alternate sequences on the polymer chain in atactic or ethylene/styrene interpolymers in which the ethylene and the sterically hindered vinylidene Representative "alternating" interpolymers are, for example, alternating

interpolymer can be described by the Bernoulli statistical model or by a first or second order Markovian statistical model, as described by J. C. Randall in Polymer Sequence interpolymers in particular, generally means that the distribution of the monomers of the do not contain more than 15 mole percent of the total amount of vinyl or vinylidene aromatic disclosure of which is incorporated herein by reference. Substantially random interpolymers Determination, Carbon-13 NMR Method, Academic Press New York, 1977, pp. 71-78, the comprising the above-mentioned monomers i), ii) and iii), and to ethylene/ styrene monomer in blocks of vinyl or vinylidene aromatic monomer of more than 3 units. The term "substantially random" as used herein in reference to the interpolymers

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degree (greater than 50 mole percent) of either isotacticity or syndiotacticity. This means peak area of the main chain methylene and methine carbons. meso diad sequences or racemic diad sequences should not exceed 75 percent of the total areas corresponding to the main chain methylene and methine carbons representing either that in the carbon-13 NMR spectrum of the substantially random interpolymer, the peak Preferably, the substantially random interpolymer is not characterized by a high

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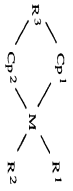
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Francis J. Timmers, both of which are incorporated herein by reference in their entirety. atmospheric up to 3000 atmospheres and temperatures from -30°C to 200°C Preferred operating conditions for such polymerization reactions are pressures from described in metallocene or constrained geometry catalysts in combination with various cocatalysts, as includes polymerizing a mixture of polymerizable monomers in the presence of one or more some amounts of homopolymer polymerization products resulting from free radical autopolymerization temperature of the respective monomers may result in formation of Polymerizations and unreacted monomer removal at temperatures above the A preferred method of preparation of the substantially random interpolymers EP-A-0,416,815 by James C. Stevens et al. and US Patent No. 5,703,187 by

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5,057,475; 5,096,867; 5,064,802; 5,132,380; 5,189,192; 5,321,106; 5,347,024; 5,350,723; interpolymers are disclosed in EP-A-514,828); as well as U.S. Patents: 5,055,438; applications are incorporated herein by reference ,374,696; 5,399,635; 5,470,993; 5,703,187; and 5,721,185, all of which patents and Examples of suitable catalysts and methods for preparing the substantially random

by the methods described in JP 07/278230 employing compounds shown by the general The substantially random lpha-olefin/vinyl aromatic interpolymers can also be prepared



groups, independently of each other; m is a group IV metal, preferably Zr or Hf, most atoms, hydrocarbon groups with carbon numbers of 1-12, alkoxyl groups, or aryloxyl substituents of these, independently of each other; R¹ and R² are hydrogen atoms, halogen wherein Cp¹ and Cp² are cyclopentadienyl groups, indenyl groups, fluorenyl groups, or preferably Zr; and R³ is an alkylene group or silanediyl group used to cross-link Cp¹ and

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by R. B. Pannell (Exxon Chemical Patents, Inc.) in WO 94/00500; and in Plastics their entirety <u>Technology</u>, p. 25 (September 1992), all of which are incorporated herein by reference in by the methods described by John G. Bradfute et al. (W. R. Grace & Co.) in WO 95/32095; The substantially random lpha-olefin/vinyl aromatic interpolymers can also be prepared

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chemical shift ranges of 43.70 - 44.25 ppm and 38.0 - 38.5 ppm. Specifically, major peaks intensities greater than three times the peak to peak noise. These signals appear in the 08/708,869 filed September 4, 1996 and WO 98/09999 both by Francis J. Timmers et al.. α -olefin/vinyl aromatic/vinyl aromatic/ α -olefin tetrad disclosed in U.S. Application No signals in the chemical shift region 43.70 - 44.25 ppm are methine carbons and the signals are observed at 44.1, 43.9, and 38.2 ppm. A proton test NMR experiment indicates that the These interpolymers contain additional signals in their carbon-13 NMR spectra with in the region 38.0 - 38.5 ppm are methylene carbons. Also suitable are the substantially random interpolymers which comprise at least one

e.g. an ethylene/styrene/styrene/ ethylene tetrad wherein the styrene monomer insertions vinyl aromatic monomer insertions preceded and followed by at least one lpha-olefin insertion, of said tetrads occur exclusively in a 1,2 (head to tail) manner. It is understood by one It is believed that these new signals are due to sequences involving two head-to-tail

monomer/ethylene tetrad will give rise to similar carbon-13 NMR peaks but with slightly and an lpha-olefin other than ethylene that the ethylene/vinyl aromatic monomer/vinyl aromatic skilled in the art that for such tetrads involving a vinyl aromatic monomer other than styrene different chemical shifts.

temperatures of from about -30°C to about 250°C in the presence of such catalysts as those represented by the formula These interpolymers can be prepared by conducting the polymerization at

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$$(ER_2)_m$$
 MR'_2 C_p

15 10 20 wherein: each Cp is independently, each occurrence, a substituted cyclopentadienyl group illustrated by the formula: cocatalyst. Particularly, suitable substituted cyclopentadienyl groups include those silicon atoms or two R' groups together can be a C₁-C₁₀ hydrocarbyl substituted 1,3to about 30, preferably from 1 to about 20, more preferably from 1 to about 10 carbon or hydrogen, halo, hydrocarbyl, hyrocarbyloxy, silahydrocarbyl, hydrocarbylsilyl containing up from 1 to about 10 carbon or silicon atoms; each R' is independently, each occurrence, hydrocarbylsilyl, containing up to about 30 preferably from 1 to about 20 more preferably Zr; each R is independently, each occurrence, hydrogen, hydrocarbyl, silahydrocarbyl, or π -bound to M; E is carbon or Si; M is a group IV metal, preferably Zr or Hf, most preferably butadiene; M is 1 or 2; and optionally, but preferably in the presence of an activating

divalent derivative of such group. Preferably, R independently each occurrence is preferably from 1 to about 10 carbon or silicon atoms or two r groups together form or hydrocarbylsilyl, containing up to about 30, preferably from 1 to about 20, more wherein each R is independently, each occurrence, hydrogen, hydrocarbyl, silahydrocarbyl, hexyl, benzyl, phenyl or silyl or (where appropriate) two such R groups are linked together (including where appropriate all isomers) hydrogen, methyl, ethyl, propyl, butyl, pentyl

tetrahydrofluorenyl, or octahydrofluorenyl. forming a fused ring system such as indenyl, fluorenyl, tetrahydroindenyl,

phenylindenyl) zirconium di-C₁-C₄ alkoxide, or any combination thereof and the like methyl-4-phenylindenyl) zirconium di-C,... alkyl, racemic-(dimethylsilanediyl)-bis-(2-methyl-4-4-phenylindenyl) zirconium 1,4-diphenyl-1,3-butadiene, racemic-(dimethylsilanediyl)-bis-(2-(2-methyl-4-phenylindenyl) zirconium dichloride, racemic-(dimethylsilanediyl)-bis-(2-methyl-Particularly preferred catalysts include, for example, racemic-(dimethylsilanediyl)-bis-

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titanium dimethyl, or any combination thereof and the like. titanium dimethyl; and ((3-iso-propyl)(1,2,3,4,5-η)-1-indenyl)(tert-butyl amido)dimethylsilane titanium dimethyl; ((3-tert-butyl)(1,2,3,4,5-η)-1-indenyl)(tert-butylamido) dimethylsilane 1-yl]silanaminato(2-)-N]titanium dimethyl; (1-indenyl)(tert-butylamido)dimethyl- silane catalysts, [N-(1,1-dimethylethyl)-1,1-dimethyl-1-[(1,2,3,4,5-η)-1,5,6,7-tetrahydro-s-indacen-It is also possible to use the following titanium-based constrained geometry

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described the influence of polymerization conditions on the copolymerization of styrene with 339) and in U.S. Patent No. 5,652,315, issued to Mitsui Toatsu Chemicals, Inc.. The described by Arai, Toshiaki and Suzuki (Polymer Preprints, Am. Chem. Soc., Div. Polym. ethylene using Me₂Si(Me₄Cp)(n-tert-butyl)TiCl₂/methylaluminoxane Ziegler-Natta catalysts. et al. (Journal of Applied Polymer Science, Vol. 53, pp. 1453 to 1460, 1994) have described ethylene-styrene copolymer. Xu and Lin (Polymer Preprints, Am. Chem. Soc., Div. Polym. pages 2387 to 2396 [1990]) and D'Anniello et al. (Journal of Applied Polymer Science, have been described in the literature. Longo and Grassi (Makromol. Chem., Volume 191, Petrochemical Industries Ltd. or US Patent No. 5,652,315 also issued to Mitsui and butene/styrene is as described in US Patent No. 5,244,996, issued to Mitsui manufacture of lpha-olefin/vinyl aromatic monomer interpolymers such as propylene/styrene Chem., Volume 38, pages 349-350, 1997; U.S. Patent No. 5,883,213 and DE-A-197 11 Copolymers of ethylene and styrene produced by bridged metallocene catalysts have been Sernetz and Mulhaupt, (Macromol. Chem. Phys., Vol. 197, pp. 1071-1083, 1997) have the copolymerization of ethylene and styrene using a TiCl₂/NdCl₂/ MgCl₂/Al(Et)₃ catalyst MgCl₂/TiCl₄/NdCl₃/ Al(iBu)₃ catalyst to give random copolymers of styrene and propylene. Lu Chem., Vol. 35, pages 686-687 [1994]) have reported copolymerization using a methylalumoxane (MAO) and cyclopentadienyltitanium trichloride (CpTiCl₃) to prepare an Volume 58, pages 1701-1706 [1995]) reported the use of a catalytic system based on Further preparative methods for the interpolymers used in the present invention

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Polymer Preprints Vol. 39, No. 1, March 1998 by Toru Aria et al. can also be employed for incorporated herein by reference. Also, although of high isotacticity and therefore not KK. All the above methods disclosed for preparing the interpolymer component are the purposes of the present invention. "substantially random", the random copolymers of ethylene and styrene as disclosed in Petrochemical Industries Ltd, or as disclosed in DE-A-197 11 339 to Denki Kagaku Kogyo

general not detrimental for the purposes of the present invention and can be tolerated. monomer at elevated temperatures. The presence of vinyl aromatic homopolymer is in aromatic homopolymer may be formed due to homopolymerization of the vinyl aromatic While preparing the substantially random interpolymer, an amount of atactic vinyl

preferably from about 45 to about 95, more preferably from about 50 to about 90 mole or cycloaliphatic vinyl or vinylidene monomer; or both; and from about 35 to about 95, percent of ethylene and/or at least one aliphatic lpha-olefin having from about 3 to about 20 percent of at least one vinyl or vinylidene aromatic monomer; or sterically hindered aliphatic preferably from about 5 to about 55, more preferably from about 10 to about 50 mole carbon atoms The substantially random interpolymer usually contains from about 5 to about 65

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and styrene and interpolymers of ethylene, styrene and at least one alpha-olefin containing from 3 to 8 carbon atoms. The most preferred substantially random interpolymers are interpolymers of ethylene

optional The presence of other polymerizable ethylenically unsaturated monomer(s) is

0.930 to about 1.040 g/cm³, most preferably from about 0.930 to about 1.030 g/cm³. The or more, preferably from about 0.930 to about 1.045 g/cm³, more preferably from about about 1.8 to about 10, more preferably from about 2 to about 5. molecular weight distribution, M_{ω}/M_{π} , is generally from about 1.5 to about 20, preferably from The density of the substantially random interpolymer is generally about 0.930 g/cm³

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aromatic homopolymers, one or more alkenyl aromatic copolymers, a blend of one or more polymers. The alkenyl aromatic polymers may be comprised solely of one or more alkenyl of each of alkenyl aromatic homopolymers and copolymers, or blends of any of the foregoing with a non-alkenyl aromatic polymer. Regardless of composition, the alkenyl Thermoplastic polymers useful in the present invention also include alkenyl aromatic

than 70 weight percent alkenyl aromatic monomeric units. Most preferably, the alkenyl aromatic polymer material comprises greater than 50 weight percent and preferably greater aromatic polymer material is comprised entirely of alkenyl aromatic monomeric units

acid, methacrylic acid, ethacrylic acid, maleic acid, itaconic acid, acrylonitrile, maleic alkenyl aromatic compounds. Examples of copolymerizable compounds include acrylic alkyl acids and esters, ionomeric derivatives, and C_4 - C_6 dienes may be copolymerized with polystyrene. Minor amounts of monoethylenically unsaturated compounds such as C_z - C_ϵ from alkenyl aromatic compounds such as styrene, alpha-methylstyrene, ethylstyrene, vinyl anhydride, methyl acrylate, ethyl acrylate, isobutyl acrylate, n-butyl acrylate, methyl having a high degree of syndiotactic configuration. A preferred alkenyl aromatic polymer is isomers of these compounds. Suitable polymers also include alkenyl aromatic polymers benzene, vinyl toluene, chlorostyrene, and bromostyrene, t-butyl styrene, including all methacrylate, vinyl acetate and butadiene. Suitable alkenyl aromatic polymers include homopolymers and copolymers derived

referred to as crystal polystyrene or polystyrene homopolymer. Science and Engineering, Vol. 16, pp. 62-71, 1989. Such polystyrene is often called also material. The term "general purpose polystyrene" is defined in the Encyclopedia of Polymer General purpose polystyrene is the most preferred alkenyl aromatic polymer

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their impact properties. Examples of suitable rubbers are homopolymers of C₄-C₆ alkyl acrylates and an interpolymerizable comonomer, especially an alkenyl aromatic or ethylidene norbornene; homopolymers of C₄-C₆ alkyl acrylates; interpolymers of C₄-C₆ aromatic monomers, and one or more C_4 - C_{ϵ} conjugated dienes; interpolymers of ethylene alkenyl aromatic polymer for use in all of the foregoing rubbery polymers is styrene. A most rubbery polymers wherein the graft polymer is an alkenyl aromatic polymer. A preferred monomer or a C,-C, alkyl methacrylate. Also included are graft polymers of the foregoing and propylene or ethylene, propylene and a nonconjugated diene, especially 1,6-hexadiene conjugated dienes, especially butadiene or isoprene; interpolymers of one or more alkenyl preferred rubbery polymer is polybutadiene or a styrene/butadiene copolymer. The monoalkenyl aromatic polymers may be suitably modified by rubbers to improve

commercially available Impact modified alkenyl aromatic polymers are well known in the art and

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polymers having a high degree of syndiotactic configuration. Suitable polymers to be employed as Component (A) also include alkenyl aromatic

rubber-modified high impact polystyrene, poly (vinyl-toluene), and poly(alphamethylstyrene). Preferred alkenyl aromatic polymers include polystyrene, syndiotactic polystyrene

10 Ų does not generate by-products in sufficient quantity to foul or coat processing equipment. subjected to melt-processing techniques adequately maintains its physical properties and lactide polymers or poly(lactide). By "melt-stable" it is meant that the lactide polymer when disposed of in an environmentally sound fashion. obtained from corn and of being biodegradable (compostable). Thus, poly(lactide) may be Poly(lactide) offers the benefits of being a renewable resource material which may be resins are supplied commercially by Cargill Dow under the designation EcoPLA the disclosure of which is incorporated herein by reference in its entirety. Suitable PLA PLA resins. Such lactide polymers are disclosed, for example, in US Patent No. 5,773,562, Lactide polymers are obtainable from lactic acid and may therefore also be referred to as Thermoplastic polymers for use in the present invention also include melt-stable

selection criteria are disclosed in US Patent No. 5,773,562 (column 14, line 35 – column 15, line 28). The poly(lactide) formulation may include a plasticizer. Suitable plasticizers and

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disclosed in US Patent No. 5,773,562 (column 7, lines 21-47). second polymer with the poly(lactide). Suitable 'second polymers' and selection criteria are To improve certain properties of poly(lactide) it may be advantageous to blend a

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polymers; polyester/polyether block polymers (e.g., HYTEL™); ethylene/propylene rubbers; styrene/isoprene/styrene (SIS) block polymers, and styrene/butadiene/styrene (SBS) block such as styrene/butadiene (SB) block copolymers, styrene/ethylene-butene/styrene (SEBS) hydrogenated conjugated diene polymer block having a hydrogenation level of greater than aromatic polymer block having a hydrogenation level of greater than 90 percent and each hydrogenated; that is the block copolymer is characterized by each hydrogenated vinyl aromatic/conjugated diene block polymers (e.g. SBS) that have been substantially and ethylene/ propylene/diene (EPDM) elastomers. Preferred elastomers are vinyl block polymers, styrene/ethylene-propylene/styrene (SEPS) block polymers, aromatic polymer blocks) to rubber polymer blocks (e.g. conjugated diene polymer blocks). 95 percent where hydrogenation converts unsaturated moieties into saturated moieties. Also, preferred block polymers have a higher ratio of rigid polymer blocks (e.g. vinyl Suitable elastomers and block polymers include, for example, block copolymers

polyhydroxyesterether or polyhydroxyether. Suitable thermoplastic phenoxy resins include polyhydroxyaminoether

to those skilled in the art. limited to, by typical grafting, hydrogenation, functionalizing, or other reactions well known The polymers used in the present invention may be modified, for example, but not

ethylenic unsaturation and at least one carbonyl group. acids, anhydrides, esters and their salts, both metallic and nonmetallic. Preferably, the Representative of compounds that contain at least one carbonyl group are the carboxylic accomplished by employing an organic compound containing at least one ethylenic Maleic anhydride is the preferred unsaturated organic compound containing at least one methyl crotonic and cinnamic acid and their anhydride, ester and salt derivatives, if any. Representative compounds include maleic, fumaric, acrylic, methacrylic, itatonic, crotonic, organic compound contains ethylenic unsaturation cponjugated with a carbonyl group. unsaturation (e.g., at least one double bond), and at least one carbonyl group (-C=O). extent, with a polar material. Graft modification of the polymers is advantageously modification renders an essentially nonpolar material compatible, at least to some limited and polypropylenes, with various unsaturated mononomers is well known in the art. Such a The graft modification of polymers, particularly polyolefins, such as polyethylenes

0.01 weight percent, and preferably at least about 0.05 weight percent, based on the other than a polyolefin, either grafted or ungrafted. one or more other polyolefins, either grafted or ungrafted, or with one or more polymers example, a graft modified ethylene-based or propylene-based polymer may be blended with polymer may be blended with one or more other polymers, either grafted or ungrafted. For compound can be grafted to the polymer by any known technique. The graft-modified more preferably it does not exceed about 2 weight percent. The unsaturated organic exceed about 10 weight percent, preferably it does not exceed about 5 weight percent, and unsaturated organic compound content can vary to convenience, but typically it does not combined weight of the polymer and the organic compound. The maximum amount of The unsaturated organic compound content of the grafted polymer is at least about

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according to established techniques. In addition or alternatively, the polymers may be chemical method, including, but not limited to, peroxide-, silane-, sulfur-, radiation-, or azide modified by suitable chain-extending or cross-linking processes using e.g. a physical or a The polymers may be sulfonated or chlorinated to provide functionalized derivatives

instance, it may be desirable to employ peroxide crosslinking agents in conjunction with are disclosed, for example, in EP-A-0 852 596, incorporated herein by reference. For heat, moisture cure and radiation steps, may be effectively employed. Dual cure systems which are herein incorporated by reference. Dual cure systems, which use a combination described in U.S. Patent No. 5,869,591 and EP-A-778,852, the entire contents of both of radiation, and sulfur-containing crosslinking agents in conjunction with silane crosslinking based cure systems. A more detailed description of the various cross-linking technologies ď

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crosslinking (crosslinked) as well as fully crosslinking (crosslinked), as long as the crosslinking results in a gel which is verifiable via ASTM D2765, Procedure A. The various based polymer. As used herein, "crosslinking" and "crosslinked" include partially polyolefin, more preferably a polyolefin as defined above, and most preferably an ethyleneor coating, the thermoplastic polymeric material or both have been cured, irradiated, or cross-linking agents can be used alone, or in combination with one another. crosslinked. Preferably, the cured, irradiated or crosslinked thermoplastic polymer is The present invention also provides a film, sheet, or coating, wherein the film, sheet,

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peroxide, and 2,5-dimethyl- 2,5-di-(t-butyl peroxy) hexyne. Dicumyl peroxide is the preferred peroxy) hexane, t-butyl-cumyl peroxide, α,α' -di(butyl peroxy)-diisopropyl benzene,di-t-butyl di-t-butyl peroxy-3,3,5-trimethylcyclohexane, dicumyl peroxide, 2,5-dimethyl-2,5-di(t-butyl agent. Additional teachings to organic peroxide cross-linking agents are seen in C.P. Park, than 120°C. For example, suitable cross-linking agents are organic peroxides, such as 1,1organic peroxides, more preferably those with one hour half lives at temperatures greater Supra, pp. 198-204, which is incorporated herein by reference Suitable heat-activated cross-linking agents include free radical initiators, preferably

or atmospheric moisture. Preferably, the silane grafted polymer is subjected to or exposed polymer backbone and thereafter subjecting or exposing the silane grafted polymer to water to water or atmospheric moisture after a shaping or fabrication operation. Alternatively, polymers may be crosslinked or cured by first grafting a silane onto the

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include those of the general formula Suitable silanes for silane crosslinking of the polymer, e.g. the ethylene polymer,

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one of the three R groups is an alkyl. independently is a hydrolyzable organic group such as an alkoxy group having from 1 to 12 lower alkyl group having 1 to 6 carbon atoms inclusive, with the proviso that not more than acetyloxy, propanoyloxy), amino or substituted amino groups (alkylamino, arylamino), or a carbon atoms (e.g. methoxy, ethoxy, butoxy), aryloxy group (e.g. phenoxy), araloxy group (e.g. benzyloxy), aliphatic acyloxy group having from 1 to 12 carbon atoms (e.g. formyloxy, when x is 1, y is 1; n is an integer from 1 to 12 inclusive, preferably 1 to 4, and each R in which R¹ is a hydrogen atom or methyl group; x and y are 0 or 1 with the proviso that

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water. The water permeating into the bulk polymer from the atmosphere or from a water the shaping or fabrication operation by reaction between the grafted silane groups and operation. However, preferably, the silane is grafted onto the polymer before shaping or commonly referred to as the "cure phase" and the process itself is commonly referred to as fabrication operations. In any case, the curing or crosslinking reaction takes place following suitable quantity of organic peroxide, either before or during a shaping or fabrication bath or "sauna". Suitable silanes may be grafted to a suitable (ethylene) polymer by the use of The phase of the process during which the crosslinks are created is

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hydrolyzable groups include methoxy, ethoxy, formyloxy, acetoxy, proprionyloxy, and alkyl present invention. Suitable silanes include unsaturated silanes that comprise an grafted onto the polymer. These silanes and their method of preparation are more fully or arylamino groups. Preferred silanes are the unsaturated alkoxy silanes which can be example, a hydrocarbyloxy, hydrocarbonyloxy, or hydrocarbylamino group. Examples of ethylenically unsaturated hydrocarbyl group, such as a vinyl, allyl, isopropenyl, butenyl, described in US Patent No. 5,266,627 to Meverden, et al.. Vinyl trimethoxy silane, vinyl cyclohexenyl or γ -(meth)acryloxy allyl group, and a hydrolyzable group, such as, for Any silane that will effectively graft to and crosslink the polymer can be used in the

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preferably the crosslinker includes vinyl triethoxy silane. the preferred silane crosslinkers for use in this invention. triethoxy silane, γ -(meth)acryloxy propyl trimethoxy silane and mixtures of these silanes are If a filler is present, then

crosslinker does not exceed 5, preferably it does not exceed 2, phr. depending several factors such as the silane itself, processing conditions, grafting maximum amount of silane crosslinker used, and typically the maximum amount of silane Considerations of convenience and economy are usually the two principal limitations on the However, typically at least 0.5, preferably at least 0.7, parts per hundred resin (phr) is used. efficiency, organic peroxide selection, the ultimate application, and similar factors The amount of silane crosslinker used in the present invention can vary widely

which is incorporated herein by reference. in the presence of a free radical initiator e.g. peroxides and azo compounds, or by ionizing The silane crosslinker is grafted to the polymer by any conventional method, typically A suitable grafting method is disclosed in WO 95/29197, the disclosure of

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20 crosslinker to initiator ratio is between 10 to 1 to 30 to 1, preferably between 18 to 1 and 24 0.10, phr. The ratio of silane crosslinker to initiator also can vary widely, but the typical initiator can vary, but it is typically present in an amount of at least 0.04, preferably at least and tert-butyl peracetate. A suitable azo compound is azobisisobutyl nitrite. The amount of peroxide, t-butyl perbenzoate, benzoyl peroxide, cumene hydroperoxide, t-butyl peroctoate, 0.06, phr. methyl ethyl ketone peroxide, 2,5-dimethyl-2,5-di(t-butyl peroxy)hexane, lauryl peroxide, compound or any one of the peroxide initiators, for example, dicumyl peroxide, di-tert-butyl But, for efficient silane grafting, organic initiators are preferred, such as Typically, the initiator does not exceed 0.15, preferably it does not exceed about

polymer, one preferred method is blending the two with the initiator in the first stage of a temperatures are typically between 160°C and 260°C, preferably between 190°C and 230°C, depending upon the residence time and the half life of the initiator. reactor extruder, such as a Buss kneader. The grafting conditions can vary, but the melt While any conventional method can be used to graft the silane crosslinker to the

copolymers, such as ethylene-vinyl silane copolymers and ethylene vinyl acetate-vinyl silane polymers Also suitable for the purpose of the present invention are moisture cure silane

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WO 01/05574

usually carried out at a dosage between about 1 megarads to about 150 megarads, gas atmosphere. Preferably, electron beam irradiation is carried out after shaping or irradiation can be carried out in air atmosphere, or in reduced oxygen atmosphere or for example 0°C to about 60°C, may also be employed. be carried out conveniently at room temperature, although higher and lower temperatures, preferably between about 3 to about 50 megarads. Further, electron beam irradiation can or 3,000,000 or 6,000,000 or higher or lower. In electron beam irradiation, the irradiation is in a range of about 50 kilovolts to about 12 megavolts with a power output capable of source can be any apparati known in the art such as an electron beam generator operating thermal heating, or any combination thereof. Electron beam irradiation is preferred. to appropriate levels which may be, for example, 100,000, 300,000, 1,000,000 or 2,000,000 supplying the desired dosage. Advantageously, electrons are employed up to 70 megarads dosages. The irradiation electrons (electron beam), ultra violet rays, X-rays, gamma rays, beta particles, controlled fabrication of the article Crosslinking by irradiation may be accomplished by the use of high energy, ionizing The voltage of the electron beam generator can be adjusted Furthermore, electron beam in inert

fabrication or processing of the polymer, but are activated by the application of temperatures (heat) substantial above normal fabrication or processing temperatures and is subsequently irradiated with electron beam irradiation at about 8 to about 20 crosslinking. ionizing energy (or both) to effectuate some measurable gelation or preferably, substantial Also, in a preferred embodiment, a polyolefin is incorporated with a pro-rad additive Suitable pro-rad additives are compounds which are not activated during normal

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organic peroxides and polyfunctional vinyl or allyl compounds such as, for example, triallyl dicumyl peroxide, di-tert-butyl peroxide, t-butyl perbenzoate, benzoyl peroxide, cumene glycol dimethacrylate, diallyl maleate, dipropargyl maleate, dipropargyl monoallyl cyanurate, compounds which have poly-functional (i.e. at least two) moieties such as C=C, C=N or combination thereof. Preferred pro-rad additives for use in the present invention are peroxy)hexane, lauryl peroxide, tert-butyl peracetate, azobisisobutyl nitrite and the like and hydroperoxide, t-butyl peroctoate, methyl ethyl ketone peroxide, 2,5-dimethyl-2,5-di(t-butyl cyanurate, triallyl isocyanurate, pentaerthritol tetramethacrylate, glutaraldehyde, ethylene Representative pro-rad additives include, but are not limited to, azo compounds,

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percent (based on the total weight of the concentrate). pro-rad additive concentration for the masterbatch is relatively high e.g., about 25 weight concentrate comprising the same or different base resin as the polymer. Preferably, the in the art. But, preferably the pro-rad additive(s) is introduced via a masterbatch At least one pro-rad additive can be introduced to the polymer by any method known

octoate, lead naphthenate, zinc caprylate, cobalt naphthenate; and the like. dioctyltinmaleate, dibutyltindiacetate, dibutyltindioctoate, stannous acetate, stannous typically between about 0.015 and about 0.035 phr. this invention. The catalyst (or mixture of catalysts) is present in a catalytic amount, carboxylate, especially dibutyltindilaurate and dioctyltinmaleate, are particularly effective for complexes or carboxylates of lead, cobalt, iron, nickel, zinc and tin. Dibutyltindilaurate bases, carboxylic acids, and organometallic compounds including organic titanates and that will provide this function can be used. Suitable catalysts generally include organic polymer. Crosslinking can also be promoted with a crosslinking catalyst, and any catalyst most preferably from about 0.015 to about 1 weight percent based on the total weight of the about 5 weight percent, more preferably from about 0.005 to about 2.5 weight percent and Preferably, the at least one pro-rad additive introduction amount is from about 0.001 Pro-rad additives are introduced to the polyolefin in any effective amount

directly downstream of a interpolymerization process. or twin screw extruder including a compounding extruder and a side-arm extruder employed mixer (for example, a Banbury mixer, a Haake mixer, a Brabender internal mixer, or a single and subsequently melt mixing in a mixer or by mixing the components together directly in a or solvents. Dry blending involves blending the individual components in solid particulate example a hydrocarbon, and combining the components followed by removing the solvent blending, that is dissolving one or both of the components in a suitable solvent, such as for the compositions by physical admixture includes dry blending, melt blending and solution immiscible or incompatible polymers can be effected by a suitable compatibilizer. Preparing formed by any convenient method. If desired or required, compatibilization between two such as blends, of thermoplastic polymers. Polymer mixtures or compositions can be The film, sheet, or coating of the invention may also comprise suitable mixtures,

substantially linear ethylene interpolymer may be formed via interpolymerization of ethylene and the desired α -olefin using a constrained geometry catalyst in at least one reactor and a The mixtures can further be formed *in-situ.* For example, blends comprising a 30

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making polymer blends by in-reactor blending using two or more catalysts in a single reactor reactor or combinations of multiple catalysts and multiple reactors. The general principle of in series or in parallel, or by in-reactor blending using two or more catalysts in a single constrained geometry catalyst or a Ziegler-type catalyst in at least one other reactor. The or combinations of multiple catalysts and multiple reactors is described in WO 93/13143, interpolymerization process is disclosed in International Application WO 94/01052 reactors can be operated sequentially or in parallel. An exemplary in-situ reference. The present polyolefin compositions can be prepared by selecting appropriate EP-A-0 619 827, and US Patent No. 3,914,362, each of which are incorporated herein by incorporated herein by reference. The blends may be prepared using two reactors operated catalyst and process conditions with a view to the final composition characteristics

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copolymer (SBS). styrene copolymer, a styrene/butadiene rubber or a styrene/butadiene/styrene block less than 30 weight percent diluent material, particularly when the diluent material is it is preferable for a polymer mixture to contain less than 50 weight percent, more preferably ethylene/methacrylate (EMA) interpolymers, and combinations thereof. In some instances, polyethylene (LDPE), EAA interpolymers, ethylene/vinyl acetate (EVA) interpolymers and and HDPE) as well as high pressure polyethylenes such as, for example, low density modified polyethylenes (for example, polybutylene and maleic anhydride grafted LLDPE Exemplary diluent materials include, for example, elastomers, rubbers and anhydride "diluent materials"), to the extent that the desired performance properties are maintained recycled and scrap materials and diluent polymers (both encompassed by the expression Thermoplastic polymers suitable for use in the present invention also include

Additives

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pigments, colorants, deodorants, fillers, plasticizers, medical ornaments such as diaper rash polyisobutylene (PIB), SANDOSTAB PEPQ™ (supplied e.g. by Ciba Specialty Chemicals), (e.g., IRGAFOS™ 168 also supplied by Ciba Specialty Chemicals), cling additives (e.g., IRGANOX™ 1010 or IRGANOX™ 1076 supplied by Ciba Specialty Chemicals), phosphites additives, including, but not limited to, antioxidants (e.g., hindered phenolics, such as ornaments, UV stabilizers, heat stabilizers, processing aid and combinations thereof Optionally, the film, sheet, or coating of the present invention may further comprise

fatty acid amides, and combinations thereof. dioxide, talc, calcium carbonate, and clay, as well as primary, secondary and substituted characteristics (slip agents) including, but not limited to, untreated and treated silicon may also contain additives to enhance antiblocking (antiblock agents) coefficient of friction Although generally not required, the film, sheet, or coating of the present invention

sheet, or coating in, for example, cushioned packaging of electronically sensitive goods. invention. Enhanced anti-static characteristics promote the usefulness of the inventive film, also be added to enhance the anti-static characteristics of the film, sheet, or coating of the combination with ethylene-acrylic acid (EAA) copolymers or other functional polymers, may Still other additives, such as quaternary ammonium compounds alone or in

Films, Sheets, Coatings

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document, a film, sheet, or coating, or a layer thereof having such fringed surface base consisting of a pattern of peaks or protrusions separated by troughs or valleys. In top item individually, e.g. "fringed film". The fringed surface microstructure may cover a desired microstructure is also referred to as "fringed film, sheet, coating or layer", including each they comprise at least one layer, which has a 'fringed' surface microstructure. In this the fringed surface microstructure. The peaks are generally non-perforated, meaning that thermoplastic polymeric material. The fringes are integral components of the layer forming The base (of the fringe layer) and the fringes (themselves) are composed of the same peaks. Such at least partially hollow peaks or so-called craters are referred to as 'fringes'. top of the peak are empty or unfilled. "At least partially hollow" includes completely hollow least partially hollow at the top, meaning that at least about 25 percent of the volume at the or cross-sectional view, the peaks have a crown-like or crater-like appearance and are at item. When magnified, the side view of a fringed surface microstructure shows a broken up part or parts, or substantially the entire fringed item, depending on the intended use of said the base at the bottom of the craters is generally not perforated. The films, sheeting, or coatings of the present invention are characterized in that

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approximate tubular or conical shapes. Various parameters can be used to characterize the diameter at the bottom of the fringe may be larger, about the same, or smaller than the like or wine glass-like shapes. The side walls of the fringe typically become thinner towards diameter at the tip of the fringe. Exemplary three-dimensional fringe structures mimic bottlegeometry of the fringes, such as the relation between different diameters. For example, the The fringes may have various three-dimensional structures. For example, they may

the top of the crater. The brim at the top of the peak or crater may be (relatively) smooth, wave-like or fuzzy. The opening may be round or elliptical.

base which is between the craters as dictated by the pattern of the base craters, followed by valleys which represent the crater holes as well as the surface of the sectional view, cutting through the center of the craters, exhibits peaks for the walls of the craters, for example a pattern of tubes or cylinders, which emerge from the base. The cross-Microscopy (SEM), the top view of the fringed surface microstructure exhibits a pattern of times or more. For example, when appropriately magnified using Scanning Electron naked eye, it is microscopically discernible, for example at an enlargement of about 10 If the fringed microstructure of the film, sheet, or coating is not discernible with the

or coating of the invention may be designed to comprise substantially similar or different index or enhanced surface area, taken alone or in any combination. The fringed film, sheet Suitable parameters include fringe density, length or height (ratios), diameters, hollowness parameters relating to the dimensions of the (overall) fringe, its (inner) hollow part, or both. fringe structures. The fringed surface microstructure can be characterized by one or more quantitative

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as microscopy or optical surface profilometry. Methods to determine the dimensional fringe parameters are known in the art, such

Celsius). Cross-sections of items with longer fringes are embedded in a medium suitable for with lengths below about 75 microns, are advantageously cut using an ultra-sharp, durable can be identified. For example, cross-sections from items with relatively short fringes, e.g sections, for example 20 to 40, should be obtained such that the hollow center of a crater embedding tissue, such as Paraplast™ wax. tool, such as a diamond knife, at very low temperatures, e.g. at -120°C (minus 120 degrees may be analyzed by optical or electron microscopy using microtoming. Several cross In brief, suitable samples, such as cross-sections, of a fringed film, sheet, or coating

from ADE Phase Shift, or VEECO Methology Group, both in Tucson, Arizona, USA. Details scanning interferometry (VSI) technology. Such profilers are commercially available, e.g. 1993, and U.S. Patents Nos. 5,133,601; 5,204,734 or 5,355,211, all by D.K. Cohen and interferometric Profiler for Smooth and Rough Surfaces", Proc. SPIE, page 2088, October on the method and the device are disclosed in an article by P.J. Caber et al., "New height variations. Optical surface profilometry is a method capable of profiling a rough surface having The method is performed with a non-contact optical profiler using vertical

when the surface is in focus. The contrast of these interference fringes(or modulation) digital processing algorithms to calculate surface heights. From these heights and the system scans downward, an interference signal for each point on the surface is increases as the sample is translated into focus, then falls as it is translated past focus. As fringes. These interference fringes are the alternating light and dark bands which appear varying heights. The source light beam is split within the interferometer. The beams characterized in that the interferometric objective moves vertically to scan the surface at fringe characterization are derivable from these structural data dimensional cross-sectional views can be generated. The dimensional parameters used for corresponding positions, a three-dimensional profile height function as well as tworecorded. The interference fringe signal is then processed (demodulated) using a series of reflected from the test surface and the reference surface recombine to form interference .P. Brophy and incorporated herein by reference. Vertical scanning interferometry is

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this technique was 0.1 nm to 1 mm standard with less than 1% error. The method provides characterize the fringe structure and determine the dimensional parameters. This method several parameters with one measurement. The method utilizes a white light source. If a two-dimensional as well as three-dimensional structural data enabling the determination of consuming sample preparation. The range of the surface height that can be profiled using has the advantage of being a non-destructive method, which does not require timebe measured by profilometry, the dimensional parameters are measured via optical fringed surface microstructure is composed of fringes with a structure which is unsuitable to microscopy. For the purpose of this invention, optical, non-contact profilometry is used to

by the lateral surface area. The arithmetic average roughness, designated as Ra in surface including peaks and valleys. The lateral surface area is the surface area measured in the surface area is the total of the exposed three-dimensional surface area being analyzed, average maximum peak-to-valley value and the average spacing of roughness peaks. microstructure data, such as the surface area ratio, the arithmetic average roughness, the reference mean plane is the three-dimensional reference surface to which all points in the metrology, is the arithmetic mean height relative to the reference mean plane. The lateral direction. The surface area ratio is calculated as the ratio of the surface area divided dataset are related. The Ra value is calculated as: The optical profilometry measurement can generate directly a number of surface

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$$Ra = \frac{1}{MN} \sum_{j=1}^{M} \sum_{i=1}^{N} |Z_{ji}|$$

fringes when the base of the film is selected as the reference mean plane. For the purpose of the present invention, the Ra values relate to the total volume of

height over the evaluation area, which is calculated as the average of each maximum peakto-valley height of the measured data array. The average maximum peak-to-valley height The average maximum peak-to-valley value is the average maximum peak-to-valley

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$$Rz = \frac{1}{N} \sum_{i=1}^{N} (Hi - Li)$$
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10 is the number of data arrays within the dataset (evaluation area). The peak-to-valley value invention, the average maximum peak-to-valley height corresponds to the fringe length or wherein Hi are the highest points and Li are the lowest points found in the data array, and N is the height difference between adjacent peaks and valleys. For the purpose of the present

preferably at least about 2000/cm². Preferably, the density is less than 10,000 per square fringes per cm². If desired, the fringe density may be selected to vary within the fringed centimeter. Most preferably, the density is in the range of from about 2000 to about 5000 surface microstructure. The density of individual fringes is at least about 1000 per square centimeter (cm²),

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profilometry. Fringe density can be determined by optical or electron microscopy, and surface

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height (H₁/2) (7). integral with the layer forming the fringes (1). (3) indicates the (total) length of a fringe (H), bottom of the fringe (D), and (6) is the (inner) diameter of the hollow center ($\mathsf{D}_{1,2}$) at half (4) indicates the depth of the hollow center or inner height (H_n) , (5) is the diameter at the Fig. 1 shows a cross-sectional view of an exemplary tubular fringe (2) which is

fringes. A particular fringe may be higher on one side. The fringes should have a minimum tip of said fringe. It is readily apparent that the height should be determined on fringes in distance between a (base) valley adjacent to the fringe whose height is determined and the height of at least about 40 micrometers (microns) or more, preferably of at least about 80 their most possible upright or vertical position, rather than on bent over or down-lying As used herein, the fringe length or height (H) is the average maximum (vertical)

fringe height is in the range of from about 200 to about 400 microns. height is less than about 1 millimeter, preferably less than 600 microns. Most preferably, the microns or more, most preferably of at least about 150 microns or more. Typically, the fringe

depth or height and diameter of the inner hollow part. These parameters can be put in relation with the overall fringe dimensions. The particular geometry of the fringes can further be characterized in terms of the

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cross-section is dissecting the center of the hollow fringes as schematically illustrated in Fig. measured from the two-dimensional cross-sectional view of the fringe structure when the determined by optical surface profilometry. These structural parameters can also be The values for $\mathsf{H}_{\scriptscriptstyle\mathsf{h}}$, D and $\mathsf{D}_{\scriptscriptstyle\mathsf{1/2}}$ can be calculated from the profile height function as

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yielding a hollow depth ratio (•) of more than 1. Preferable, • is below 1.3, more preferably = H_n/H). The depth of the hollow center (H_n) may exceed the (outer) height (H) of the fringe, average depth of the hollow center (H_n) to the average (maximum) height of the fringe (H) (• The hollow depth ratio (•) is defined as the ratio of the average inner height or

the vertical portion of the fringe starts. While the ratio may be higher than 1, fringe center at half (inner) height ($D_{1/2}$) and the diameter at the bottom of the fringe (D) (• = $D_{1/2}/D$). a hollow diameter ratio of at least about 0.1 or higher. In case the hollow diameter ratio is characteristic of the fringed surface microstructure of the present invention preferably have structures with values of about 1 or lower are preferred. At least partially hollow craters higher than 1, the measurements should be determined by optical microscopy The diameter at the bottom of the fringe (D) is determined at the point of inflection at which The hollow diameter ratio (•) is defined as the ratio of the diameter of the hollow

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the hollow depth ratio (*) with the hollow diameter ratio (*) (* = * \times * \times 100 = $H_n/H \times D_{1/2}/D \times D_{1/2}$ unfilled volume at the top of the fringe. The Hollowness Index (•) is calculated by multiplying Hollowness Index (•). The Hollowness Index is indicative of the degree of hollowness, or the preferably, more preferably the Index is 100 or lower, most preferably about 90 or lower. For more preferably 40 percent or higher. The Hollowness Index may be more than 100 (e.g., in 100). The Hollowness Index (*) is typically about 15 or higher, preferably about 25 or higher, case (•) is higher than 1). The Hollowness Index should be below 260, preferably below 130 The hollow depth ratio (•) and the hollow diameter ratio (•) are used to calculate the

are considered as being completely hollow. the purpose of the present invention, fringes having a Hollowness Index of 100 or higher

fringe (having the same outside geometric dimension). according to the present invention has a greater surface area than a corresponding solid profilometry. Generally and as evident from Fig. 1, an at least partially hollow fringe the density of the fringes. area which depends on the fringe height (H), the hollowness, the diameters (D and $\mathsf{D}_{\scriptscriptstyle{1/2}}$) and Another parameter suitable to characterize the fringe structure is the total surface The Surface Area Ratio can be obtained by optical surface

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preferably between about 1 and 3. The aspect ratio (A) is the ratio of the fringe height (H) and the fringe diameter (D) (A =H/D). Preferably, the fringes have an aspect ratio of between about 1 and about 5, most Another parameter useful to characterize the fringe structure is the aspect ratio (A).

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from about 100 to about 300 microns. center distance between two adjacent fringes. Preferably, the center-to-center distance is Another parameter characterizing the fringed surface microstructure is the center-to-

thicknesses, i.e. areas of reduced thickness (valleys or troughs) and areas of greater characterized by a distinct surface texture consisting in a pattern of different (surface) thickness (peaks or protrusions). Advantageously, this pattern is predetermined and may be starting from a suitable precursor film, sheet, or coating. The structure of said precursor is one layer of the film, sheet, or coating according to the present invention may be obtained irregular or regular. The particular three-dimensional fringed surface configuration characterizing at least

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according to the present invention will be referred to as "precursor" (film, sheet, or coating). pattern of different surface thicknesses and suitable to give a fringed surface microstructure microbubbles of the foam. In the following, a film, sheet, or coating characterized by such having a foamed surface layer. In such case, the precursor protrusions are formed by the This includes a film, sheet, or coating which in part is characterized by such pattern of different surface thicknesses Alternatively, the precursor may be a foamed structure, e.g. a film, sheet, or coating

treated such that the protrusions are essentially longitudinally extended. Preferably, such To obtain the fringed surface microstructure the precursor surface is mechanically

using mechanical means, such as a mechanical pulling force and/or an abrasive device. longitudinal extension or stretching of the precursor protrusions is the result of treatment

sheet or coating is pulled off the matrix surface under certain conditions process and subsequent formation of the fringed surface microstructure when the film, precursor film, sheet, or coating on the matrix surface in a continuous compression molding involves a matrix surface suitable to create a surface texture and comprises formation of the fringed surface microstructure occur in a single continuous process. Such in-line process performed in an in-line process, meaning that formation of the precursor structure and the The mechanical treatment of the precursor film, sheet, or coating is preferably

pressure. The negative structure may, for example, consist of very fine cavities. The cavities precursor such that the thermoplastic material closely contacts said reverse structure under surface temperature of the moving belt or roller is adjustable. depth), shape and entry angle (with respect to the matrix surface). Advantageously, the may have various geometries - primary variables include cavity dimensions (diameter, negative or reverse approximation with respect to the desired surface texture of the The matrix surface, for example a moving belt or a roller surface, presents a

at a certain angle. Appropriate control and defined conditions for the mechanical treatment film, sheet, or coating from the matrix surface by exerting on the protrusions a tractive force are essential to make a fringed surface microstructure according to the present invention. extension of the precursor protrusion occurs during the removal or peel-off of the precursor In such a continuous process, mechanical treatment resulting in the longitudinal

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an abrading device should affect, for example, the fringe length and/or the texture of the surface characteristics and properties. The additional mechanical deformation by means example with an abrasive material, is optional, but may be desired to enhance fringed Mechanical post-treatment of the thus obtained fringed film, sheet, or coating, for

surfaces may have different, similar or equal surface roughness and shape. The surfaces pair of rollers, preferentially enabling temperature control, with defined surface qualities. The one surface, referred to as matrix surface, is characterized by the presence of numerous preferably, be installed for a secondary (separate) compression molding process. At least fringed film, sheet, or coating comprises a set of surfaces, as presented for example by a Suitable equipment for such continuous compression molding process to make a part of the primary equipment used to make the (base) film, sheet, or coating, or,

suitable material, or preferably by eroding techniques (electrical, chemical, lasering). polymeric or natural particles, or fibrous materials, by mechanical or chemical treatment of a fringe number or density. Such matrix surface structure can be provided by a porous natural, metallic, polymeric), by sintering of a suitable material, such as metallic, ceramic, material, an open cell, foamed material, by woven or entangled fibrous structures (e.g., about 100 microns. The number or density of cavities should correspond to the desired Preferably, the cavities are substantially regular or symmetrical. cavities with a projected area of at least about 1000 square microns and a depth of at least

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surface is a rubber, advantageously having a Shore A hardness in the range of about 70 to ceramic, e.g. on an appropriate support, such as steel. A particularly preferred matrix such as a steel core, a polymer, e.g. coated on an appropriate support, such as a steel, or a about 85, preferably a halogen-elastomer, such as a fluoroelastomer. Advantageously, the matrix surface is lasered with fine cavities having or approximating the form of cylinders. Preferred matrix surfaces are steel, a rubber, e.g. covering an appropriate support,

the matrix surface is in the range of from about 45 degrees to about 90 degrees, preferably are known in the art. The matrix surface is applied on a device suitable for processing a film, cylinder-like cavities, the angle of incline of the axis of symmetry of the cylinder relative to sheet, or coating, such as a roller or a belt. Typically, for symmetrical cavities, such as Suitable techniques and technologies to make the matrix surface and the cavities

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finished product which is not facing the matrix surface) may be accomplished in the form of a film, sheet, or coating. If desired, lamination (to that surface of the semipolymer solution, film, sheet, or coating is applied on the matrix surface. Advantageously polymer mass, e.g. in the form of a polymer melt, polymer dispersion, polymer suspension, penetration of the polymer mass into the cavities is facilitated by heating the surface of the used to force the polymer mass into the cavities of the matrix surface. Preferably, surfaces suitable to apply pressure onto the polymer mass, such as a roller or a belt, is simultaneously with the continuous compression molding step. One or more counter and preferably, the polymer mass is applied in form of a semi-finished product, in particular select the appropriate temperature. The surface of the polymer mass is molded such that forming the protrusions. Generally, the person of ordinary skill in the art is readily able to mass to a temperature which is close to, preferably above the melting point of the polymer To prepare the desired precursor having a pattern of different surface thicknesses

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the presence of additional sources of energy for curing or crosslinking after the surface and/or curing and/or cross-linking may be performed on the matrix roller or belt, optionally in fed into the compression molding equipment is a suspension or emulsion (additional) drying temperature which is about at or advantageously below the Vicat point. If the polymer mass or coating is cooled such that the thermoplastic material forming the fringes has a when the film, sheet or coating is stripped off the matrix roll. Advantageously, the film, sheet off the matrix surface. Most preferably, thermoplastic material is in the semi-molten state solidified, when the fringe-forming force is applied and the film, sheet or coating is stripped sources, such as an air knife or cooling water. Preferably, the thermoplastic material is not achieved by appropriately cooling the precursor, if desired using additional external cooling Low(er) viscosity of the polymer is preferred. After the surface molding, the film, sheet, or pressure and temperature (of the polymer mass surface and the relevant equipment). shaped according to the structure of the counter surface of the pressuring device. The molding has occurred (tensile) strength to pull the molded elevations out of the cavities. Sufficient strength is coating is released or pulled off from the matrix surface, which step requires sufficient pressure roll and the matrix roll. Key parameters during formation of the precursor are roll surfaces of the precursor reflect the surface characteristics of surfaces, e.g. both rolls, the a suitable precursor structure. At the same time, the other surface of the polymer mass is individual, distinct surface elevations or protrusions are formed in the cavities, thus yielding

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tractive force is dependent on adhesion or interaction between the polymer comprising the fringe layer and the matrix roll surface and the release angle. protrusions involves subjecting the precursor and the protrusions to a tractive force. The microstructure. Thus the mechanical treatment of the precursor providing deformation of the characterizing the precursor surface are elongated to give the fringed surface During the release process of the precursor from the matrix surface, the protrusions

the angle between the fringed film, sheet, or coating during the release process and the preferably at least about 90 degrees. The release angle should be less than about 170 greater than about 20 degrees, more preferably at least about 45 degrees and most through the point of release. The release angle should be greater than 10, preferably the release angle is the angle between the fringed film, sheet, or coating and the tangent no more covered with the film, sheet, or coating. In case the matrix surface is (on) a roller, matrix surface. In this context matrix surface means that part of the device which is free and A crucial parameter in this step of mechanical treatment is the release angle, that is

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further extend the protrusions of the precursor, thus still further increasing the surface area Release angle, take-off speed and polymer surface temperature are selected such as temperature and below the crystalline melting point of the polymer forming the protrusions. surface. The temperature at the polymer surface should be above the glass transition degrees. The release angle is impacted by the angle of the cavities in relation to the in respect to the precursor structure.

arrangement of the rolls may be vertical or horizontal. pressure in the nip can be measured according to methods known in the art. The linear compression distance at the center (axis) of the rolls is the negative gap. The counter (or preheat) roll and the matrix roll at the temperature and pressure applied. The roll, forming a nip. The nip width depends upon the compression of either or both of the formation. In the preferred process, the matrix roll is in contact with a pre-heat or counter comprises a (cavity) filling step and a release or peel-off step, both of which affect fringe invention is a roller-based continuous compression molding process. Such process A preferred process to make the fringed film, sheet or coating of the present

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suitable to make the fringed film, sheet or coating of the invention. Fig. 2 schematically shows an example of a calander with vertically roll arrangement)

increased or reduced by means of one or more idler rolls (12) which set the point of initial depending on the desired polymer temperature in the nip, the contact distance can be coating contacts the pre-heat roll (11) for a predetermined distance. If appropriate, and above the melting point of the polymer forming the fringed surface. The base film, sheet or (relatively) smooth surface. Advantageously, the temperature of the preheat roll (11) is at or sheet or coating desired to be converted into a fringed surface microstructure have a pressure in the nip can be measured according to methods known in the art. of the preheat roll and the matrix roll (13) at the temperature and pressure applied. The contact on the preheat roll. The nip width depends upon the compression of either or both roll (11) representing the counter surface. Preferably, at least those parts of the base film, So-called base film, sheet or coating is fed from a feeder roll (10) onto a pre-heat

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polymer layer coming in contact with the matrix roll (13) preferably is at or above the melt heating of the surface of the film, sheet or coating. The temperature of the particular speed, and any additional heat that may be supplied via auxiliary methods, such as infrared the nip will be dependent upon the distance that the film wraps the preheat roll and the line The actual amount of preheat time that the film, sheet or coating experiences prior to

pressure in the nip. Advantageously, these variables are measured and/or controlled using temperature should be such that the majority to all of the polymer is molten. Critical from or peel-off of the matrix surface (release step) advantageously involves a take off step yields the precursor which is then cooled as it travels along the matrix roll. Release (13) should be negative. Another critical parameter is the interaction of the polymer and the roll axis should be at least about 1 MPa, or higher. The gap between the two rolls (11) and Maximum nip pressure measured at the center of the nip width in line with the center of the conventional methods known in the art. For example, nip contact pressure can be formation include polymer type, and (its response to) line speed, temperature in the nip, and variables in the 'filling step', i.e. in the filling of the matrix cavities, which affects fringe temperature of the polymer when it leaves the nip contact width. For blends, the the matrix roll surface and the polymer, and their temperatures), and the release or peel or to increase production speed. The amount of force/tension required to peel the film, may also be present. Such device serves to facilitate removal of the film from the matrix roll, supporting cooling of the film, sheet or coating on the matrix roll, such as an air knife (14, system, for example comprising a chill roll (15) and a winder (16). Optionally, a device matrix roll surface. The polymer should have sufficient wettability to the matrix roll. The filling measured using commercially available pressure measuring film, e.g. Fuji Prescale Film. angle. Fig. 3 illustrates the determination of the peet angle (B): adhesion of the polymer to the matrix roll surface (which is a function of the composition of sheet or coating from the matrix roll for a given fringe layer polymer is dependent upon the

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adhesive peel (or controlled adhesive failure) between the polymer and the matrix surface obtained, while ensuring that the polymer is completely removed from the cavity due to an precursor so that a maximum amount of draw or elongation of the fringe layer polymer is fringe formation, it is important to optimize the adhesion; line speed and the strength of the This results in hollow fringe formation with optimized fringed length. The greater the adhesion, typically the higher the resulting peel angle. For hollow

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desirable to use a sacrificial backing layer or protective layer. The sacrificial layer can be, oriented or non-oriented PLA (polylactic acid) film, or natural fiber substrates, and/or a but not limited to, oriented or non-oriented PET film, aluminum foil, non-woven/fleece oriented or non-oriented polypropylene film, oriented or non-oriented polystyrene film, the equipment used in making said item or to facilitate release of the item, it may be 'release aid', such as PTFE, silicon, etc. as equipment coating materials (e.g. on the In order to avoid sticking of the film, sheet, or coating according to the invention to 48-

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fringed layer material are suitable as sacrificial layers. acids, etc.). Film or sheet exhibiting a melting point which is at least 20°C higher than the pressure roll or counter roll), or plasticizer or migratory additives in the polymer matrix (fatty

polymer is favored by employing a thermoplastic having a low viscosity at the relevant requirements of the envisaged end-use applications. Optimal filling of the cavities with the conditions for making the precursor and the fringed surface and the specific performance dimensionally stable, resilient and resistant to deformation under load at ambient the fringed surface microstructure is selected such that the fringes (after formation) are processing temperature temperature. Generally, polymer selection will be dictated by the particular process Advantageously and preferably, the thermoplastic polymer or polymer blend forming

in the range of about 0.900 g/cm³ to about 0.960 g/cm³ (e.g. DOWLEX™ or ATTANE™ copolymers available from The Dow Chemical Company), or polymers produced via high heterogeneous ethylene-based polymers, such as Ziegler/Natta polymers having a density ethylene-based polyolefins, including homogeneous ethylene-based polymers and process combining optimal filling in the continuous compression molding processs and pressure processes. mechanical treatment by application of a tractive force are polyolefins more preferably Preferred polymers for use in the fringed layer obtainable by the above-described

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polymer surface temperature, and hardness (difference). Advantageously, the treatment fabrics, such as a fleece or a non-woven having the required surface hardness. Suitable not limited to, sandpaper or sand-paper type materials, brushes, steel wool, or abrasive parameters include deforming speed, applied pressure, temperature, particularly the the type of abrasion material and the particular technique of treatment therewith. Critical is readily apparent to the person skilled in the art that the length of the fringes is affected by surface coating than the polymer surface to be treated. Suitable abrasives include, but are treatment with an abrasive device. Preferably, the abrasive device has a harder surface mechanical post-treatment, such as treatment relying on the principle of friction, for example and the tips may become frayed. Best results are obtained by circular, oscillating or linear such that the fringed surface structure is enhanced, e.g. the fringe structure is elongated materials for the brush are, for example, nylon, glass-fiber or metal wire, such as bronze. It movements, or any combination thereof. The general direction of these movements relative Optionally, the fringed film, sheet, or coating thus obtained may be subjected to

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the surface. For each combination of surface hardnesses, i.e. surface hardness of the rotational or oscillating treatment, sequential treatment steps can improve the final quality of at least two consecutive steps of either unidirectional or angled treatment. In the case of movement and speed to obtain a particular fringed surface microstructure, e.g. fringe length abrasion material and surface hardness of the polymer surface, exists an optimum type of any combination thereof. If the treatment is essentially unidirectional, it is preferred to have to the fringed surface and surface orientation may be longitudinal, transverse, angled, or be frayed, torn and elongated. magnified, e.g. using the SEM technique, the tops of the walls of the craters may be seen to and Hollowness Index. Such optimization is within routine experimentation. When

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compression (for example, molding), roll milling or calendering, or any combination thereof. solvent, solvent including an aqueous medium), extrusion (for example, blown or casting), any known process including, but not limited to, solvent casting (for example, dispersion in a combination thereof. One or more films may be laminated to form a multi-layer structure Extrusion as used herein is intended to include co-extrusion, extrusion coating or any Also, a film may be laminated to a woven or non-woven fabric forming a composite A film fed into the above described compression molding process can be made

blowing agent, to make a foamed film. To make foamed film structures or layers, either about 2.0 mm. The foams may be crosslinked or non-crosslinked the foams typically are from about 0.01 mm to about 5.0 mm, preferably from about 0.02 to the art. The foams may be open or closed cell, according to ASTM D2856. The cell sizes of foam densities from 1 g/cm³ to as low as 0.01 g/cm³. Suitable blowing agents are known in physical or chemical blowing agents may be used to achieve suitable foam densities, e.g One or more layers of the film may be expanded, for example with a conventional

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casting, extrusion, roll milling, compression or calandering, or any combination thereof. A sheet can be made by any known process including, but not limited to, solvent

coating or involving a roller and doctor blading. A coating can be made according to methods known in the art, e.g. by extrusion

surface microstructure include soft, velvety and textile-like feel or touch, matte appearance. Performance attributes which can be specifically provided or enhanced by the fringed fringed film, sheet or coating particularly benefit from the fringed surface microstructure. Fringed films, sheets, or coatings, and articles of manufacture comprising such

may be made to provide additional functional properties, such as breathability, increased and/or of a proper fringe structure. If desired or required, the fringed film, sheet or coating attributes, for example by selection of a proper polymer material or polymer materials designed and optimized to specifically exploit or favor one or more of these performance imprintability, controlled release properties, storage capabilities, and protective properties liquid repellency, anti-skid and good grip properties, surface variability and surface heat resistance, or barrier properties. Depending on the intended end-use application the fringed surface microstructure can be

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and for household applications. for industrial, personal or medical hygiene applications, for apparel or protective clothing, packaging or as packaging material, for use as labels or signs, in automotive applications, manufacture are useful, for example, for decoration or as decorative materials, for performance attributes afforded by the fringed surface microstructure such articles comprising a fringed film, sheet or coating. Based on the specific advantageous The present invention also relates to an article of manufacture made from

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plastic articles of manufacture used in soft touch, soft feel applications. For example, soft to manufacture plastic based articles for which such improved haptics are desirable, e.g. sheet, or coating of the present invention has improved haptics and is very pleasant and invention affords an excellent soft, velvety and textile-like feel or touch, The fringed film, with (direct) skin contact, for example, furniture covers including outdoor furnishing, such as or coatings according to the invention are also useful to make decorative fabrics or articles door panels, seat covers, headliners, and steering wheel covers. The fringed films, sheets, automotive interior surfaces, including but not limited to instrument panels, console liners, touch, soft feel materials are desirable in the automotive industry, for example on easy on a person's skin. Based on this property such fringed film, sheet, or coating is useful desk covers, cushions, including patio air cushions, drapes, bedspreads, and table cloths; protective gear; such as medical drapes and gowns, athletic clothing, raincoats, hats and ears or wings, cuffs, side panels, and back sheets; (disposable) garments apparel, and condoms, incontinence articles, such as diapers, or parts thereof including e.g., top sheet, medicinal and personal hygienic articles, including for example bandages, band aids ostomy bags; mattresses including air mattresses; life vests; artificial leather; floor covering; shoes; gloves, and protective gear, artificial skin, skin imitations, toys, industrial hygienic caps; grip bands; (bag) handles; linings, for example for luggage, bags or handbags, or More specifically, the fringed surface microstructure characterizing the articles of the

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feel. Such articles can be useful as scouring or abrading devices, or to provide anti-skid in differentiated haptics and tacticity, affording a (relatively) rough feel rather than a soft end-use applications requiring a soft feel the fringe length is advantageously at least about advantageous to obtain a soft feel. The use of polymers having a higher modulus will result 150 microns or more. Generally, use of polymers having a low modulus polymer will be articles, such as wipes or disposable toilet seat covers, head rest covers and the like. For

cling sings, merchandise bags, and projection and movie screens. linings of jewelry boxes and luggage; anti-glare surfaces such as billboards and window decorative overlays, including wall, floor, or ceiling covering products; artificial flowers; relying on excellent aesthetics and matte optical appearance including, for example The films, sheets, or coatings according to the invention are useful to manufacture articles reduced gloss (relative to the precursor). Gloss is determined according to ASTM D-2457. film, sheet, or coating according to the invention is a matte (mat) appearance and further Another benefit provided by the fringed surface microstructure characterizing the

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differentiated surface tension, as determined by the test methods according to ASTM Dfeminine hygiene top sheet, (disposable) table cloth and shower curtains. liquid repellant surfaces, e.g. rain gear, such as hats, coats, or ponchos, protective clothing, management such that water beads up and drains off, is required, such as clothing with invention are useful for applications and articles where surface water or liquid drain 2578 or DIN 53364, and water contact angle. The fringed films, sheets, or coatings of the Another advantageous property characterizing the fringed film, sheet, or coating is

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as reflected e.g. in the dynamic coefficient of friction, which affords anti-slip or anti-skid and manufacture with a surface made from or comprising a fringed film, sheet, or coating of the better grip performance attributes to a fringed film, sheet, or coating, or an article of surface. The fringed surface microstructure provided herein typically provides an increased example, the same or a different fringed surface, or a glass, metal, ceramic or polyolefin determined according to ISO 8295, relative to various surfaces of interest, including, for invention. The dynamic coefficient of friction of a particular fringed surface may be coefficient of friction as compared to non-fringed (plain) film of the same composition. The useful for a variety of applications and articles including, but not limited to floor cover better grip and anti-slip properties make the fringed film, sheet, or coating of the invention Furthermore, the fringed surface microstructure also provides a frictional behavior,

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closing systems, e.g. VELCRO TM -like systems, serving trays (e.g. as used in air planes or shower mats, roofing, tarps for construction, truck bed covers, box coatings, self-adhering backings, furniture covers, gloves, table wear, bathroom articles, such as shower shoes restaurants), grip bands or tapes, and non-skid operation room table cover.

wearer's ability to hold onto slippery or wet instruments or improve the feel against a another material. For example, the coating may serve to enhance grippage, increase the Prior to the curing or leaching steps, the gloves may be dipped into a bath for coating with difficult to accomplish, or by peeling the latex off the form while inverting it at the same time. The gloves may be removed from the form by blasting them off by air, which is rather cured any undesirable solvents or materials are allowed to leach out during a leaching step. latex is applied to the forms by dipping the forms into the latex bath. Before the product is forms are heated and a coagulant for the latex is added to support curing of the latex. The dispersion, such as latex. The hand forms are cleaned, rinsed and dried. Subsequently, the structure. Thus the secondary dipping step of the conventional process becomes In such process, the forms are employed as matrix surface. The peel/inversion process to invention can advantageously be used to make gloves with a fringed surface microstructure. person's skin. The continuous compression molding process according to the present remove the gloves from the form is applied such as to give the fringed surface micro-The process for making gloves comprises dipping hand forms into a suitable

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in the art, including for example adhesive tape peel tests, abrasion tests, or simply visual of ink adhesion, color density (vividness), print definition employing suitable methods known fringed surface may be (im)printed with or without ink. Printability may be evaluated in terms particularly surface printability or imprintability (embossing), as reflected in the possibility to create distinct areas of different topography, e.g. using suitable impression techniques. The film may be reverse printed, e.g. by printing the fringed microstructure and viewing from the dependent on the angle of view thus creating a unique 3D-type appearance. Clear fringed inks and should be selected based on the fringe layer polymer. sides of the fringe. Acceptable ink adhesion is possible for water based and solvent based appearance. The ink may reside inside and/or between the fringes as well as at the tips or Another benefit afforded by the fringed surface microstructure is surface variability, Color density may vary

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surface structure is marked, it may also be noticeable by a person with reduced visual ability or a blind person. Nevertheless, color or ink may be added to enhance the effects optic effects are achieved by the resulting difference in surface structure and appearance the fringes in a selected regular or irregular pattern to form a smooth surface, haptic and/or Thus printing effects may be obtained without the need for color or ink. If the difference Using, for example, a hot stamp, hot air or friction to re-melt and/or compress part of

60°C to about 160°C, more preferably from about 70°C to about 120°C. The image or polyethylene are in the range of from about 50°C to about 200°C, preferably from about thermoplastic polymer forming the fringes may be employed. Suitable temperatures for printing or sealing. For example, a stencil or print negative which is a metal or thermoset which does not involve contacting the fringed surface, e.g. by applying a jet of hot effluent that a three-dimensional effect is obtained. The effects are also obtainable in a process pattern of the stencil or print negative is applied by pressing it on the fringed surface such material that can be heated without distortion to temperatures above the melting point of the film, sheet, or coating 'a priori'. create areas of differentiated topography resulting in print-like images. Alternatively, to imprinting effects is to abrade the fringes on the surface by a mechanical process and thus (e.g. air) similar to an air brushing printing process. Another process suitable to obtain obtain the described effects, the fringed surface may be created only on parts of the basic The desired imprinting effects may be realized relying on the same principles used in

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painted surfaces, e.g. in an automobile, such as dash boards, instrument panels etc., period of time. The advantage of improved scratch resistance also applies to printed or scratch and abrasion resistant their particular properties will be maintained for a longer such as barrier coatings, or metallization. By making the (barrier) coatings or deposits more item. The benefit of enhanced abrasion and scratch resistance is also afforded to coatings, scratched or abraded, thus enhancing the durability and appearance of the fringed article or metallized films, e.g. used in the packaging of food, medical items or electronics, and barrier The fringed surface microstructure can be printed with ink and resist ink from being

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of the present invention is particularly useful to make novelty balloons, book covers wrapping paper, floor covering, ceiling covering or wall covering, labels, including, for Based on its surface variability and/or imprintability the fringed film, sheet, or coating

coatings, e.g. for drinking cups, greeting cards, and party articles. example, in-mold labels and stretch labels, e.g. for bottles, bill boards, or paperboard

filtration media, diaper top sheets, e.g. coated with diaper rash ointments, anti-static sheets, of antibiotics and may be useful e.g. in wound covers or shower curtains, the controlled numerous applications and articles, including but not limited to articles comprising a outside surface, an inside surface, as well as the surface between the hollow tubes. The surface area (e.g. as compared to a filled protrusion). The hollow tube structure has an and poultry diapers. Craters which are at least partially hollow provide a greatly enhanced anti-fog films, and for HF weldability, thermal insulation, sound deadening, meat packaging storage properties can also be exploited in wipes, tissues, carrier substrates for catalysis, release of fragrances or the controlled release of drugs. Enhanced carrying, capturing or controlled release system. For example, such system may provide for the controlled release affords enhanced carrying, capturing or storing properties, which can be exploited for porous membrane may be put over the top of the reservoirs to allow the soft surface to packages for cosmetics or perfumes, scented wall covering, or as filtration systems. A charge on the surface further enhances the ability to pick up substances, such as dirt or creams with the soft side against the skin. Antibacterial agents can be placed on the can be placed in the reservoirs to maintain moistness for burn wounds or for diaper rash from a porous material the drug is delivered through the base polymer. Emollients or salves or EVOH resin, is coated over the top of the reservoirs and the base of the film is made breath to the skin (drugs) or air (fragrances). If a barrier, for example made from SARANTM useful e.g. in transdermal drug delivery systems, fragrance delivery systems, such as soft controlled release of substances, such as drugs or fragrances. Such reservoir structures are coated onto. Such reservoir structures allow, for example, the gradual diffusion or the surface to prevent mold and mildew formation, e.g. in shower curtains. Placing a static are suitable reservoir structures, which other substances can be embedded into The increased surface area of the fringed film, sheet, or coating of the invention

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suitable absorbent fillers include, for example, superabsorbents used in hygiene incorporated into the polymer (as a type of filler) before the fringed film, sheet, or coating applications, talc and calcium carbonate made, for example in order to enhance the absorption of coatings or ink. Examples of Furthermore, absorbent materials can be coated onto the fringed surface or be

invention in apparel, such as protective clothing, or in hygienic articles, such as (diaper) performance attribute desired or required for use of the fringed film, sheet, or coating of the extruded on top of an inherently breathable monolithic layer. Water vapor breathability is a microporous layer or the fringed film can be stretched. A (thin) fringed layer can also be polyamide. A (thin) fringed layer may be coextruded on top of a previously stretched thermoplastic elastomer or another highly water vapor transmitting polymer, such as stretched microporous films, or monolithic films, for example made from a thermoplastic polyurethane, a copolyesterether thermoplastic elastomer, a copolyesteramide (moisture) vapor permeable or breathable. For example, breathable films include filled and The fringed film, sheet, or coating according to the invention may be made to be

which breathability is desirable, including athletic wear, side panels in diapers or adult range of 1000 to 6500 g/m²/day. Such film is suitable for apparel or other applications for having a high WVTR. The WVTR measured at 38°C and at 90% relative humidity (using polymer). The films may be mono-layer or multi-layer structures, preferably each layer (e.g., PEBAX ™ copolymer), and thermoplastic polyurethane (e.g., PELLETHANE™ make (inherently) breathable fringed film include, for example, polyether block copolymer water repellency performance or liquid impermeability. Thermoplastic polymers suitable to different function in the composite, such as noise reduction or elastic recovery. breathable monolithic fringed film may be laminated to another material which provides soft touch has been provided by laminating a non-woven to the film. If desirable, the vapor transmission rate (WVTR) and has a soft touch. In conventional breathable films, the incontinence products ASTM method E96) should be in the range of 500 to 10,000 g/m²/day, more typically in the Advantageously, the fringed surface microstructure is designed so as to provide the desired For example, a breathable monolithic fringed film is characterized by a high water

or coating comprising poly(lactide) are particularly suitable for disposable articles e.g. boutique bags, and protective covers, e.g. for cars, bikes or boats. Fringed film, sheet, include pouches, e.g. pouches for flowable materials, such milk or detergent pouches, bags, against scratching, or cushioning, and/or printability. Exemplary packaging applications packaging applications, e.g. packaging applications requiring enhanced protection, e.g. The fringed films, sheets, or coatings according to the invention are useful for

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with other modifying elastic or non-elastomeric materials are also contemplated being useful with a styrene content of less than 40 weight percent. Blends of these polymers alone or in the present invention. The low modulus of elastic films is also advantageous in terms of copolymers, specifically at a density less than 0.89 g/cc, and ethylene-styrene interpolymers rubbers, ethylene-propylene rubbers, thermoplastic polyurethanes, ethylene-alpha olefin and B is an elastomeric midblock derived, for example, from conjugated dienes or lower alkenes), chlorinated elastomers and rubbers, ethylene propylene diene monomer (EDPM) copolymers (where A is a thermoplastic endblock such as, for example, a styrenic moiety permanent set. Elastic polymeric materials include, for example, AB and ABA block or graft point at which the elastic material begins to pull a load is designated as the percent subsequently released to the original position before stretch, and then stretched again. The film. Permanent set is the converse of elasticity. A film is stretched to a certain point and strain (doubled the length). Recovery refers to a contraction of a stretched material upon initial sample length)]x 100. Elasticity can also be described by the "permanent set" of the expressed as: [(maximum stretch length--final sample length)/(maximum stretch length-termination of a force following stretching of the material. Percent recovery may be about 50 percent of its stretched length after the first pull and after the fourth pull to 100% preferred embodiment, the elastic film, sheet or coating of the invention achieves at least original form upon release of any pressure or force applied to the film material. In comprises a material which is highly stretchable and which reverts to its original or nearly The fringed film, sheet or coating may be elastic. The elastic film of the invention

may be oriented according to methods known in the art. In a post-treatment step, the fringed film, sheet or coating according to the invention

increasing the wetting tension. The greater the polar component the more actively will the for example, corona treatment. Corona treatment increases the polarity of the surface, thus surface react with different polar interfaces. characteristics of a fringed film may be modified by techniques known in the art, including, If desired, the fringed film may be further treated (post treatment) and the surface

or articles of manufacture comprising such film, sheet, or coating, characterized by a fringed parts of the surface, or the entire surface. Such mono-layer structure is preferably made surface microstructure on one side, or on both sides. The fringed microstructure may cover One aspect of the present invention relates to a mono-layer film, sheet, or coating,

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polymer blend may be composed of two homogeneously branched ethylene-based preferably comprising the polymers indicated as being preferred. For example, a suitable polypropylene polymer, or a (poly)lactide, or a mixture or blend of thermoplastic polymers, branched ethylene polymer, and a substantially random ethylene/styrene interpolymer, a based polymer, including for example a heterogeneously or, preferably, a homogeneously from a suitable thermoplastic polymeric material indicated as being preferred herein-above. The mono-layer may be made from a single thermoplastic polymer, preferably an ethylene-

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steering wheel covers, or articles with much exposure to the sun, such as outdoor furniture applications in the automotive industry, such as automotive interior applications, such as or coating affords the benefit of enhanced heat resistance, as required, for example, by using the methods and techniques described in more detail herein-above. Such film, sheet, polymer is a polyolefin, most preferably an ethylene (inter)polymer. Crosslinking is achieved surface microstructure. the thermoplastic polymeric material or both have been cured, irradiated or cross-linked. coating comprising a thermoplastic polymeric material, wherein the film, sheet, or coating Advantageously, curing, irradiation or crosslinking is performed after forming of the fringed The present invention further provides a mono-layer or multi-layer film, sheet, or Preferably, the cured, irradiated or cross-linked thermoplastic

one layer has a fringed surface microstructure and is comprised of a thermoplastic polymer sheet or coating wherein the fringed layer is an outer layer are preferred layer or layers. The fringed layer may be an inner and/or, an outer layer. A multi-layer film, as defined herein. The fringed microstructure may be present on one or both sides of said Generally, in a multi-layer film, sheet, or coating according to the invention, at least

on the intended end use application, cost considerations, and the like characteristic to the overall structure. The composition of these layers is chosen depending In a multi-layer structure, each layer will serve a particular function or provide some

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properties, thermal properties, optic properties sealing characteristics, chemical resistance, referred to as a tie layer. If a barrier layer is desired or required for the intended end use mechanical properties, or abuse resistance. An adhesion promoting interlayer is also characteristics, e.g. add bulk to the structure, promote interlayer adhesion, provide barrier For example, layers may serve to provide particular structural or functional

application, it is selected so as to meet the targeted degree of gas or moisture (im)permeability.

polyethylene terephtalate (PET), oriented polypropylene (OPP), ethylene/vinyl acetate nylon, ethylene/vinyl alcohol (EVOH) copolymers, polyvinylidene chloride (PVDC), anhydride grafted polyethylene. (EMAA) copolymers, LLDPE, HDPE, LDPE, graft adhesive polymers, such as maleic (EVA) copolymers, ethylene/acrylic acid (EAA) copolymers, ethylene/methacrylic acid or more than one layer in the same film structure. Suitable materials include, for example, Various materials can be used for these layers, with some of them being used in one

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10 EVOH copolymer or SARAN™. For example, a gas barrier layer may be made from vinylidene chloride copolymer,

LLDPE, ULDPE, VLDPE, POP, EVA copolymer, EAA copolymer and ionomers Examples of thermoplastic polymers appropriate for use in the seal layer include

on cost considerations and include recycled materials. Representative polymers are, for HDPE, polypropylene polymers and blends thereof. example, LLDPE, such as ethylene/butene copolymers, LDPE, EVA copolymer, (recycled) Thermoplastic polymers for use in the bulk layer are advantageously chosen based

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seven layers. Preferred are multi-layer films having three or more layers fringed multi-layer structures according to the invention typically consist of from two to advantageous mechanical properties include, for example, ethylene/C₄-C₈ copolymers. Examples of thermoplastic polymers appropriate for use in a layer providing

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Permeability may be provided by a porous membrane.

wherein the respective compositions of the layers are chosen from thermoplastic polymer film, sheet, or coating including the following general two-layer and three-layer structures resins which provide the requisite functional properties: For example, it is within the scope of the present invention to provide a multi-layer

Fringed/Seal
Fringed/Adhesion
Fringed/Bulk/Seal
Fringed/Bulk/Adhesion
Fringed/Mechanical/Seal

Fringed/Bulk

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Fringed/Mechanical/Adhesion Permeability/Fringed/Barrier

added to promote inner layer adhesion or add bulk, as appropriate For each of these general structures, further internal or external layers may be

feminine hygiene articles, bandages and wound cover materials, and incontinence articles, preferably a film, in a hygienic product, including a disposable hygienic product, such as for example diapers. The present invention also relates to the use of a fringed film, sheet or coating

geomembranes The present invention also relates to the use of fringed film, sheet or coating

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fringed film, particularly a multi-layer coextruded fringed film. The fringed surface ostomy bags, urological collection bags and other liquid containment pouches polyolefin or polyester nonwovens, to provide a laminate structure that can provide patient bags are examples of patient-worn bags. In many commercial applications today, barrier microstructure is particularly advantageous and desirable for bags that are worn by a urological collection bags and other liquid containment bags or pouches comprising a of a separate nonwoven textile from a laminated bag structure. "nonwoven" or textile-like feel to a coextruded barrier film structure can result in elimination comfort when worn in contact with the body. The use of fringed film to impart soft films are combined with soft nonwoven textiles, such as spunbonded or meltblown person and are in contact with his or her skin. Ostomy bags and urological collection leg Furthermore, the invention relates to medicinal collection bags, such as ostomy bags, Furthermore, the present invention relates to the use of a fringed film in medical

Chemical or MXD6® nylon from Mitsubishi Gas Chemical Company), polyester (PET and (PVDC, such as SARAN® resin from The Dow Chemical Company), ethylene-vinyl alcohol can be achieved by use of conventional barrier polymers such as polyvinylidene chloride that provide good oxygen, carbon dioxide and odor barrier properties. Barrier properties producing a coextruded barrier film, it is desired to provide outermost film layers that provide (such as BLOX® thermoplastic epoxy resins from The Dow Chemical Company). PETG, such as EASTAR® polyester resins from Eastman Chemical Company) or phenoxy (such as GRILON® or GRILAMIDE® resin from EMS-Chemie, CAPRON® nylon from Allied (EVOH, such as EVAL® resin from EVAL Company of America), nylon or amorphous nylon Of particular interest for incorporation of a fringed film are coextruded barrier films

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10 15 ÚΊ the polyolefin polymer. and maielc anhydride (MAH) grafted polyolefins such as PE-graft-MAH, PP-graft-MAH and Suitable ethylene copolymers include EVA, EMA, EAA, ethylene-n-butyl acrylate (EnBA), such as ethylene copolymers with 1 to about 40 weight percent of polar comonomer. EVA-graft-MAH. These resins typically have 0.05 -- 1.4 weight percent MAH grafted onto together. Tie layers may include polyciefin copolymers that are polar monomer modified, butylene-styrene (SEBS). These resins typically have poor affinity for and adhesion to the styrenic block copolymers such as styrene-butadiene-styrene (SBS) and styrene-ethylenevarious barrier resins, necessitating the use of sultable tie layer resins to bond the layers copolymer (coPP), chlorinated polyethylene (CPE), thermoplastic polyurethane (TPU), ethylene-acrylic acid (EAA), and polypropylene homopolymer (PP) or polypropylene Interpolymers (ESI), ethylene-vinyl acetate (EVA), ethylene methyl acrylate (EMA), polyethylene (HDPE), metallocene or plastomer polyethylene (mPE), ethylene-styrene strength and sealability, such as with polyolefin resins. Sultable polyolefin resins include low density polyethylene (LDPE), linear low density polyethylene (LLDPE), high density

typical barrier film will have a barrier layer of at least about 0.2 mils (5 μm) in thickness. unacceptable bulging or protrusion of the bag from under the patient's clothing, eliminating the discreteness of wearing the appliance. In order to achieve good barrier properties, a patlents body. This results in uncomfortable wearing of the bag, as well as aesthetically bags or collection bags that are stiff and inflexible, resulting in poor conformation to a mils (250 μm) in thickness, and preferably about 2.0 mils (50 μm) to about 6.0 mils (150 μm) in thickness. Thicknesses significantly greater than the preferred range can result is ostomy Barrier films of the present invention are typically about 1.0 mils (25 μm) to about 10

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"A" skin layers of LLDPE, mPE or coPP at 35% each of the overall gauge. This can be asymmetrical film structure can be represented as ABCBD, wherein the "D" layer is represented as a 35/10/10/10/35 layer ratio of the 4.0 mil (100 μ m) film. A possible 5-layer "B" adhesive tie layers of EVA or EVA-g-MAH at 10% each of overall gauge; and outermost film might have a "C" barrier layer of PVDC or EVOH or PETG at 10% of the overall gauge; barrier layer, adjacent adhesive layers on either side of the barrier layer, and outermost skin tle or bonding layers, and the "A" layers are skin layers. A representative 4.0 mil (100 μm) as a "ABCBA" structure, wherein the "C" layer is a barrier layer, the "B" layers are adhesive layers on each side of the film. Such multi-layer structure can symbolically be represented Suitable film structures include, for example, a symmetrical five-layer film with a core

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"B" adhesive layer can be used to thermally bond or seal two plies of the film together, such adhesive tie or bonding layers, and the "A" layer is a skin layer. In this case, the exposed can be represented as "ABCB" wherein the "C" layer is a barrier layer, the "B" layers are that is useful in making ostomy or collection bags is a 4-layer asymmetrical structure. different skin material than the "A" material. Another representative barrier film structure as during heat sealing of a bag seam.

the direct contact of a plastic film against the skin. Direct contact of a smooth film can permeation of the excreted human waste products through the pouch, also prevent properties of the containment bag or pouch, which are required to eliminate odor bacteria or infectious growth between the bag and skin of the patient. The barrier due to abrasion against the skin in addition to hot and moist contact, and can promote cause perspiration and "clinging" of the film to the skin, can cause blisters and sores to of woven and nonwoven laminates or coverings over the bag is commonly used to eliminate against the body, it is desired that they exhibit excellent haptics, softness and comfort. Use appliance is worn. Use of a one or two fringed layers according to the present invention as moisture in the form of skin perspiration from escaping from the region where the pouch or wearing a pouch desires the appliance not to make any noise or rustling sound while especially preferred for coextruded layers on an ostomy or collection bag. These films provide a softer, more conformable surface. Elastomeric films, such as those made from modulus and low durometer "hardness" be used for the outermost layers of the film so as to escape from behind the appliance. It is further desired that films with low flexural or tensile skin layer or layers 'A' provides the desired soft skin contact and can allow perspiration to moving around, as commonly occurs when conventional plastic films are flexed, folded or promote quietness, which a desired attribute of a bag or pouch appliance. A patient TPU, CPE, styrenic block copolymers, ESI and homogeneous polyolefin resins, are For ostomy bags, urological collection bags and other appliances which are

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provide suitable bonding via thermal welding, impulse sealing, high frequency (also known fittings may be made of polyolefins such as polyethylene, EVA or EMA, or of TPU or flexible usually made of a material compatible with the film and bondable to the film. Tubing or on the film utilized for medical bags, pouches or appliances. These fittings and tubing are vinyl (PVC). Film structure and surface compositions may need to be formulated so as to radio frequency) sealing or mechanical fastening. Additionally, use of liquid curable It is additionally desired to utilize plastic tubing and/or plastic fittings or connectors

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adhesives or hot melt applied adhesives can be utilized to adhere tubing or fittings to a bag

embodiment, the floor or wall covering is a multi-layer film or sheet comprising at least one product comprising a fringed film, sheet or coating according to the invention. In a preferred print layer and/or the backing layer, the fringed microstructure facing the floor. Used as a covering product, the fringed layer may be present as the surface or top layer, the print print layer, an optional reinforcement layer and/or an optional foam backing layer. In a wall layer having a fringed surface microstructure. The product may further comprise an optional the fringed layer affords anti-slip properties. If present, the reinforcement layer is an be imprinted, if desired. If the fringed layer serves as a print layer, the fringed microstructure surface layer, the fringed microstructure is designed to afford soft touch and feel and may layer or both. In a floor covering product, the fringed layer may be the surface layer, the further contain appropriate additives, e.g. flame retardant additives to insure compliance which can be in excess of 4 meters in width. The optional intermediate reinforcement layer intermediate layer providing dimensional stability to the product, preferably a floor covering, and preferably covered (on the top) by a transparent upper wear layer. As a backing layer with flame retardant requirements and other regulatory requirements. polymer, a melt processed polyolefin polymer or a latex composition. The product may Suitable resilient foam backing layers comprise or consist of a solvent dispersed polyolefin insulation characteristics to the product as well as provide a barrier against microorganisms. resilient cushioning properties to the product and can also impart noise and thermal material or a non-woven polymeric material. If present, the foam backing layer imparts Preferably, the optionally intermediate reinforcement layer is a non-woven glass fleece may be a melt processed polyolefin polymer or a non-woven or woven textile material. is designed to provide proper surface variability and imprintability. This layer is imprinted In yet another aspect, the present invention relates to a floor or wall covering

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xamples

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as limiting the scope thereof in any manner. The following abbreviations are used: ESI are indicated in MegaPascal (MPa) and the temperatures are in degree Celsius. substantially random ethylene/styrene interpolymer; MFI or MI = melt flow index (measured kg/190°C according to ASTM D-1238, condition E); N/T = not tested; the pressures The following Examples are illustrative of the invention, but are not to be construed

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with 0.8 cm of Viton™ rubber. The rubber is laser engraved with a regular pattern (13) with a total diameter of 11.76 cm (10.16 cm diameter steel roll covered on each side calender comprises a steel pre-heat roll (11) with a diameter of 12.7 cm and a matrix roll Figure 2 shows a schematic drawing of the calender used to make the fringed Example 1: Preparation and Characterization of Fringed Films 1A – 1D

(10) is 100 microns thick and has the following general structure: For Fringed Films 1A and 1B, the (basic) smooth film fed into the calender via the feeder roll

of about 110 microns and a depth of about 270 microns. The center to center distances are

185 microns (closest neighbor) and 267 microns (diagonal neighbor).

of cylindrical dead-end holes (cavities, 2860 per square centimeter) which have a diameter

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50% Outer Layer, 30 % Core Layer, 20 % Backing Layer.

0 15 25 layer consists of 92 % HDPE (0.29 MI, 0.9605 g/cc) and 8% TiO₂ Masterbatch. The process ethylene/octene copolymer (30 Mt, 0.902 g/cc) and 6% of a Slip Masterbatch containing 5% ethylene/octene copolymer (30 MI, 0.885 g/cc), 27% of a homogeneous substantially linear layer, is made from a blend consisting of 67% of a homogeneous substantially linear The outer layer which comes in contact with the matrix roll and is converted into the fringed gap of about 3 - 3.5 MPa (= 30 - 35 bar), contact nip width of 12 mm. Line speed is 1 erucamide in 6 MI, 0.900 g/cc polyolefin plastomer. The core layer consists of 87% LDPE fluid pressure measured at the inlet to the pistons which apply pressure to the axis of the given peel angle. The temperature of the chill roll (15) is 15° C. Hydraulic Pressure is the is run with a 21 mils (= 533 microns) negative gap width (applied maximum pressure in the (0.8 MI, 0.924 g/cc) and 13% TiO₂ Masterbatch (60% TiO₂ in LDPE or LLDPE). The backing matrix roll. In this particular configuration the matrix roll presses against the pressure m/min. The winder speed is equal to line speed plus the necessary delta to maintain the temperature is measured at angle $(\alpha) = 90$ degrees (see Fig. 3). (preheat) roll. Film off the matrix roll is measured near point P₁ in Fig.3. Matrix roll

30 on the equipment variables a (= 4 inches), b (= 2 inches), c (= 1.38 inches), and d (= 2.315Fig. 3 shows the parameters used in the calculation of the release angle (β) which is based between the tangent to rubber roll at the point of detachment and the film. The angle is inches) as defined in Fig. 3, and on the wrap angle of the matrix roll (13). β is the angle based on the rules of analytic geometry.

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the point of detachment from the matrix roll (13), P_2 is the point of touch to the chill roll (15), calculation detailed below, the release angle calculates as 112.3. L is the length of the segment connecting P_{τ} to $P_{z},\,\beta$ is the angle between the tangent to the matrix roll (13) at the point of detachment P, and segment L. Based on the method of In Fig. 3, O, is the point of origin of the circle C_1 , O_2 is the point of origin of the circle C_2 , P_1 is

The calculation of release angle is based on Fig. 3:

The length of the line segment L is given by,

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$$L = \left[\left[a + c + d \cdot (1 - \cos \alpha) \right]^2 + \left(d \cdot \sin \alpha - b \right)^2 - a^2 \right]^{0.5}$$

origin P_1 . Equation(2) describes the circle C_2 with radius a and origin O_2 . P_{2} , (X,Y) have to be calculated. Equation(1) below describes a circle with radius L and Replacing the origin of a rectangular coordinate system at O₁, the coordinates of the point

$$L^2 = (x + d \cdot \sin \alpha)^2 + (y - d \cdot \cos \alpha)^2$$

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$$a^2 = (x+b)^2 + [y - (d+c+a)]^2$$

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(1) and (2). equivalent to finding the points of intersection between the circles described by Equations The values (X,Y) are obtained by solving Equations (1) and (2) simultaneously. This is

25 The tangent to circle C₁ at P₁ is:

$$m_1 = \tan \alpha$$

The slope of the line segment L is

$$m_2 = \frac{Y - d \cdot \cos \alpha}{X + d \cdot \sin \alpha}$$

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The angle between L and the tangent to C_1 at P_1 is

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$$\beta = \tan^{-1} \left(\frac{m_2 - m_1}{m_1 \cdot m_2 + 1} \right)$$

equipment variables and wrap angle. For •=60, • is calculated as 112.3 degrees proper angle corresponding to • is chosen by considering the geometry for given sets of Because the intersection of the two lines gives rise to two complementary angles, the

prevent the desired abrupt peeling. Furthermore, any bending moment in the film may modulus of the film is low. Sticky fringes may reduce the actual peel angle because they reduce the peel angle. The following table correlates wrap angle and release or peel angle. The calculation of the release angle assumes that the peeling is abrupt and the flexural

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Wrap Angle	Calculated Peel Angle
0	25
15	45
30	68
45	90
60	112
75	133
90	151

sampling area is 460 x 600 square microns. WYKO NT3300 Optical Profiler and the Vertical Scanning Inteferometry (VSI) mode. The The dimensional parameters characterizing the fringed microstructure are measured using a 10

15 The following process conditions are used to produce Fringed Film 1A:

Wrap Angle: 75 - 90°

Peel Angle: 133 - 151°

Hydraulic Pressure (MPa): 17

Preheat distance on pressure roll: 4.5 cm

Film Temperature into gap: 98°C

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Film Temperature off Matrix Roll: 51°C

Matrix Roll Surface Temperature: 61°C

Pressure Roll Surface Temperature: 126°C

Air Knife Delta: 20°C

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of preheat is dependent upon the distance and the line speed. Air knife delta is defined as Preheat distance is the distance that the film is in contact with the preheat roll. The amount the reduction in the matrix roll surface temperature as achieved by applying surface air to the back of the matrix roll as the process is running.

are as follows: The fringe characteristics of Fringed Film 1A, as determined by optical surface profilometry,

Fringe height (H): 237 microns

Fringe Diameter at base (D): 155 microns

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Surface Area Ratio: 9.6

Hollow Depth Ratio (•): 1.1

Hollow Diameter Ratio (•): 0.52

Hollowness Index (•): 57

Aspect Ratio: 1.5

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The following process conditions are used to produce Fringed Film 1B:

Wrap Angle: 60-75°

Peel Angle: 112-133°

15 Hydraulic Pressure (MPa): 17.1

Preheat distance on pressure roll: 22.5 cm

Film Temperature into gap: 91°C

Film Temperature off Matrix Roll: 46°C

Matrix Roll Surface Temperature: 59°C

20 Pressure Roll Surface Temperature: 115°C

Air Knife Delta: 13°C

of preheat is dependent upon the distance and the line speed. Air knife delta is defined as the reduction in the matrix roll surface temperature as achieved by applying surface air to Preheat distance is the distance that the film is in contact with the preheat roll. The amount

the back of the matrix roll as the process is running.

are as follows: The fringe characteristics of Fringed Film 1B, as determined by optical surface profilometry,

Fringe height (H): 220 microns
Fringe Diameter at base (D): 175 microns

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Surface Area Ratio: 5.7

Hollow Depth Ratio (•): 0.5

Hollow Diameter Ratio (•): 0.35

Hollowness Index (*): 18

Aspect Ratio: 1.3

used to make Fringed Film 1C: A smooth film (with a thickness of 100 microns) of the following structure and composition is

U 50% Outer Layer, 30 % Core Layer, 20 % Backing Layer.

g/cc polyolefin plastomer. The core layer consists of LDPE (0.7 MI, 0.922 g/cc). The backing ethylene/octene copolymer (30 MI, 0.902 g/cc), 20% of an ethylene/vinyl acetate copolymer ethylene/octene copolymer (30 MI, 0.885 g/cc), 22% of a homogeneous substantially linear layer consists of HDPE (0.3 MI, 0.947 g/cc). (8.0 Ml, 12% VA) and 6% of a Slip Masterbatch containing 5% erucamide in 6 Ml, 0.900 layer, is made from a blend consisting of 52 % of a homogeneous substantially linear The outer layer which comes in contact with the matrix roll and is converted into the fringed

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m/min. The winder speed is equal to line speed plus the necessary delta to maintain the pressure in the gap of about 2.5 MPa (= 25 bar), contact nip width of 9 mm). Line speed is 1 given peel angle. The temperature of the chill roll is 20° C. The process is run with a negative gap width of 12 mils (= 300 microns) (applied maximum

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The following process conditions are used to produce Fringed Film 1C:

Wrap Angle: 15°

Peel Angle: 45°

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Hydraulic Pressure (MPa): 16.5

Preheat distance on pressure roll: 22.5

Film Temperature into gap: 91°C

Film Temperature off Matrix Roll: 75°C

Matrix Roll Surface Temperature: 75°C

Pressure Roll Surface Temperature: 117°C

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The fringe characteristics of Fringed Film 1C, as determined by optical surface profilometry,

are as follows:

30 Fringe height (H): 335 microns

Surface Area Ratio: 17.4

Hollow Depth Ratio (•): 1.0

Hollow Diameter Ratio (•): 0.76

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Hollowness Index (•): 76

Aspect Ratio: 2.8

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in long(er) fringes with a high(er) Hollowness Index Fringed Film 1C shows that the presence of a polar comonomer in the fringed layer results

used to make Fringed Film 1D: A smooth film (with a thickness of 100 microns) of the following structure and composition is

50% Outer Layer, 30 % Core Layer, 20 % Backing Layer.

styrene comonomer (25 MI). The core layer consists of LDPE (0.7 MI, 0.922 g/cc). The layer, is made from a substantially random ethylene/styrene interpolymer containing 30% of backing layer consists of HDPE (0.3 MI, 0.947 g/cc). The outer layer which comes in contact with the matrix roll and is converted into the fringed

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is 1 m/min, the chill roll temperature is 20°C. pressure in the gap of about 2.5 MPa (= 25 bar), contact nip width of 9 mm). The line speed The process is run with a negative gap width of 12 mils (= 300 microns) (applied maximum

The following process conditions are used to produce Fringed Film 1D:

Wrap Angle: 30°

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Peel Angle: 68°

Hydraulic Pressure (MPa): 17.6

Preheat distance on pressure roll: 4.5 cm

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Film Temperature into gap: 49°C

Film Temperature off Matrix Roll: 54°C

Matrix Roll Surface Temperature: 55°C

25 are as follows: The fringe characteristics of Fringed Film 1D, as determined by optical surface profilometry,

Fringe height (H): 174 microns

Surface Area Ratio: 3.65

Hollow Depth Ratio (*): 0.48

Hollow Diameter Ratio (*): 0.55

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Hollowness Index (*): 26

Aspect Ratio: 1.5

<u>Example 2:</u> Scanning Electron Microscopy (SEM) of a Fringed Film

density of 0.912 g/cc and a MFI of 1.0 g/10 min (ATTANE™ SL 4101 resin available from White Masterbatch. The outside layer is made from 94 weight percent of a LLDPE having a MFI of 0.7 g/10 min(LDPE 300R available from The Dow Chemical Company) and 15 wt.% core layer is made from 85 weight percent of a LDPE having a density of 0.923 g/cc and a ethylene/octene copolymer having a density of 0.902 g/cc and a MFI of 30 g/10 min. The density of 0.885 g/cc and a MFI of 30 g/10 min and 67 weight percent of a homogeneous masterbatch, 27 weight percent of a homogeneous ethylene/ octene copolymer having a the fringed surface microstructure, is made from a blend of 6 weight percent of slip The Dow Chemical Company), and 6 weight percent of White Masterbatch. is evaluated using Scanning Electron Microscopy. The fringed layer, namely the layer with A three-layer fringed film (layer ratios: 60% fringed layer / 20% core / 20% outside)

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examined and photographed using a Hitachi S-4100 FEG SEM. All photographs are taken at 2.0 kV accelerating voltage. Angstroms of chromium using a Denton Hi-Res 100 chromium sputtering system, then conductive carbon tape and carbon paint. The sample is coated with approximately 60 magnification. A portion of the fringed film is cut and mounted on an aluminum sample stub with Photographs are taken at 50x, 250x, and 500x

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exhibits peaks for the walls of the tubes or craters, followed by valleys which exhibit the base film. The cross-sectional view of the film, cutting through the center of the tubes, tubes can also be described as craters. The walls of the craters or tubes are raised from the see into the tube as it is open on top but closed at the bottom by the base film. Such hollow pattern or matrix of hollow tubes or cylinders that emerge from the base film. One is able to as dictated by the base pattern. crater holes themselves as well as the surface of the base film which is between the craters When magnified using the SEM technique, the top view surface of the film exhibits a

<u>Example 3:</u> Fringed Film treated by Electron Beam Crosslinking

crosslinking. The film specimens are placed in cardboard fileholders and irradiated using to the beam multiple times to give total dosage levels of 160, 288 and 448 kGray (16, 28.8, The radiation dose per pass is 32 kGrays (3.2 Mrad). Three fringed specimens are exposed 12 MeV, 10 kW electron beam source at Sterigenics International, Inc. (San Diego, CA). Specimens of the fringed film of Example 2 are treated via electron beam

determined by xylene extraction per ASTM D2765 method, Procedure A with two 44.8 Mrad), respectively. The gel content (insoluble fraction) of the crosslinked films is the calculation of gel content. The results of the gel testing are as follows: sample holders are weighed. dried in the oven for 12 hours under at least 28 in Hg vacuum. After an hour of cooling the 12 hours and are then transferred to a vacuum oven preheated to 150°C. The samples are (w/v) 2-2'-methylene -bis-(4-methyl-6-tertiary butyl phenol)). Samples are kept in xylene for round bottom flask containing xylene boiling in reflux condition and an antioxidant (0.85% made of stainless steel wire cloth. The sample holder is then immersed into a three neck The ASTM D-2769 method involves placing about 0.3 g of sample into a sample holder exceptions. The films are not conditioned and are tested as received without grinding The difference in weight before and after extraction is used in

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Sample	Radiation Dose, kGy	%Gel
		(Average)
Sample 3-A	160	20.3
Sample 3-B	288	48.6
Sample 3-C	448	54.6

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80°, 90° and 110° Celsius each for 10 minutes. After being cooled to room temperature, the samples are mounted on glass slides, then placed on a hot plate and exposed to 60°, 70°, hot plate to evaluate the ability of the various samples to maintain their surface texture. temperature the surface of the tubes or craters is very sharp or vivid. The edges of the ability of the fringed surface observed at room temperature to remain intact. At room films are evaluated using the SEM technique described in Example 2 to determine the appear to overlap or touch the surfaces of the other tubes (Condition 0). As the surface frayed tops are very thin and wispy in nature. The tops of some of the tubes or craters shrink back to the domain of their tube (Condition 1.0) and then eventually there is no becomes affected by rising temperature the frayed, thin, wispy tops first begin to become craters (Condition 3). Eventually the diameter of the crater becomes much smaller as do the distance between the tubes or craters becomes smaller as do the distances between the more regular and oval and appear to be thicker and not at all frayed or wispy and the overlapping of the wispy tops (2.0). With increasing heat the tops of the surfaces become more isolated with less overlapping to other craters indicating that they are beginning to Samples 3-A, 3-C, and a non-crosslinked control are also exposed to heat using a

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the definition of the craters is considerably less discernable than Condition 3 but still distances between the craters as the wall shrinks closer to the base of the film. At this point various samples are provided below. observable (Condition 4). The final view is when the craters have completely shrunk back into the base of the film and are no longer discernable (Condition 5). The rating of the

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Condition 4	Condition 3	Condition 5	110° C
Condition 2	Condition 1	Condition 2	90° C
Z	Condition 1	Condition 2	80° C
N/T	Condition 0	Condition 1	70° C
N/T	Condition 0	Condition 1	60° C
N/T	ארו	Condition 0	25° C
Sample 3-A	Sample 3-C	Control	Temperature

prevent the surface from completely melting at the 110°C, with the highest level of Celsius for the non-crosslinked control. The lowest and highest level of crosslinking both postpone the loss of the frayed, wispy appearance until 80° Celsius as compared to the 60° crosslinking keeping it most intact. As observed by SEM the highest level of crosslinking (Sample 3-C) appears to

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upper service temperature of the films. Applications benefiting from enhanced temperature cushion covers (capable of withstanding high temperatures in Arizona sun or attic panels, instrument panels, dash boards and headliners, as well as outdoor furniture resistance include, for example, automotive interior applications such as fabric or vinyl door This example demonstrates the ability of cross-linked fringed films to increase the

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Example 4: Printing Tests

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on a conveyor belt and when they pass an electronic eye the lot number or product type is Printing Ink by Marsh Called UN 1210 FD. The FD indicates fast drying. The bags move by polyethylene heavy duty shipping sacks. The ink used is a methyl ethyl ketone based non-contact printing apparatus, a Marsh LCP ML4 Ink Jet System which is used to print printed on the side of the bag in a macro dot matrix form. The tested versions are a non-I. The film of Example 2 is evaluated for printability. The first printing test involves

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the side of a bag which is run past the electronic eye. composition of the film of Example 2. Approximately 4 inch x 4 inch samples are taped to textured control sample and a fringed film sample, each having the structure and

finish and the dots do not at all appear raised but sunk into the film. It has a softer one can see the raised ink dots which sit on the top of the film. The fringed film has a mat Printed Appearance: the printed appearance of the control film is very glossy and

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smudging. wiping a finger across the surface of each specimen. None of the specimens exhibits Smudge/Smear Test: within 30 seconds the bags are tested for smudging by lightly

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from the films. A qualitative analysis gives the following results: adhesion. Strips of adhesive tape approximately 0.5 inch \times 2 inch are pressed with moderate but consistent pressure across the surface of each film and are then peeled away Ink Adhesion: immediately following the smudge test, the bags are evaluated

fringed film: approximately 10-20% of each ink dot is removed non-textured control film: approximately 60-75 percent of each ink dot is removed;

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the non-textured film with the tape applied only at moderate pressures. The level of ink to the fringed sample the removal of ink is still approximately half of what is observed with textured sample. Even after applying the adhesive tape with dramatically higher pressures what is observed immediately after the printing. adhesion to the films after a week of sitting at room temperature remains consistent with The fringed sample exhibits significantly improved ink adhesion relative to the non-

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dot in a back and forth motion until it appears that no change in the color intensity of the dot and scratch tests. The abrasion test involves wiping a pencil eraser head across a single ability of the ink to remain in place. The results of these two tests are described below. is observed by additional wiping. The scratch test involves scraping a moderately sharp implement such as a coin or a fingernail across the printed film surface and observing the Ink Abrasion/ Scratchability: the specimens are also evaluated by simple abrasion

Sample	Abrasion Test	Scratch Test
Non-Textured Film	Within 5 strokes of the	Ink is easily scraped off by
	eraser the surface ink dot	scraping a sharp
	is removed and only a	implement across the
	light stain remained	surface. Entire dots can
	below.	be removed after
		repeated scrapings with a
		light stain remaining
		below.
Fringed Film	After 25 strokes of the	Single swipes across the
	eraser the ink dot is still	dots reveal no obvious
	not removed from the film	change in the ink dots.
	surface and the remaining	Repeated swipes are able
	dot looks significantly	to lighten the dots but do
	darker than the light stain	not approach ink removal
	of the non-textured film	achieved by the eraser.

50% fringed/ 30% core/ 20% outside) of the following composition: II. Another printing test involves evaluation of a three-layer fringed film (layer ratios:

copolymer having a density of 0.885 g/cc and a MFI of 30 g/min, 67 wt.% of a homogeneous ethylene/octene copolymer having a density of 0.902 g/cc and a MFI of 30); core layer: 87 wt.% LDPE 300R, 13 wt.% White Masterbatch; fringed layer: 6 wt.% slip masterbatch, 27 wt.% of a homogeneous ethylene/octene

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outside layer: 92 wt.% ATTANE SL 4101, 8 wt.% White Masterbatch.

paper items. A smooth (non-textured) base film of the corresponding structure and composition is used as control film. The fringed film is evaluated on an ink jet desktop printer typically used for printing

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definition of the dot pattern. In addition, it has a very matte finish and appears to have the appearance is blurred. The fringed film does not bead up and shows significantly better dimensionality of the film surface appears to create a visual three-dimensional appearance appearance of a soft-colored pencil look as opposed to a typical ink link. The physical multiwetted well outside of the boundary of the dot pattern common to ink jet printers. The multidimensional effect can benefit various applications for which the (visual and haptic) depth or creating a better shading effect by alternating colors. Such soft shading effect or as well. This effect provides an enhanced and improved visual appearance giving a greater including, for example, doll skins and automotive seats. depth appearance of a textile in combination with the cleanability of a plastic are desired Appearance of the films: on the non-textured film ink beads up on the surface and

smudge beyond the boundary of the printed object. This finding could indicate that the ink of the film. The ink on the fringed film comes off on the finger, but does not appear to textured film smudges beyond the boundaries of the printed object and smears onto the rest on the fringed is more firmly encapsulated in the hollow tubes of the film. both the non-textured control and the fringed film is not completely dry. The ink on the non-Smudge test: a finger smudge test is conducted after 12 hours revealing that ink on

recessed centers of the craters. that the ink resides on the tips of the fringe, in between the fringe, as well as in the Inspection by microscopy of the top view of the ink jet printed fringed surface reveals

Example 5: Pouch comprising a multi-layer fringed film

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adhesive tie layer, and a ULDPE-LDPE blend innermost skin layer (ABCBD structure). polyethylene copolymer, an EVA adhesive tie layer, a PETG core barrier layer, an EVA film line. The 5.0 mil (125 μm) thick film is comprised of a outermost layer of homogeneous outermost layer is 50% (2.5 mil, 62 μm) of the film gauge and consists of 96% Affinity™ of 100% Elvax™ 3190 (DuPont, 25% vinyl acetate, 0.95 g/cc, 2.0 g/10 MFI). The core resin). The two EVA tie layers are each 10% (0.5 mil, 12 μm) of the film gauge and consist antiblock concentrate (20% silicon dioxide, 3.5% stearamide, 3.5% erucylamide in EVA SM1300 (The Dow Chemical Company, 0.902 g/cc, 30 g/10 MFI) and 4% CN4420 slip / 60% Attane™ 4201 ULDPE (The Dow Chemical Company, 0.912 g/cc, 1.0 g/10 minute melt The innermost skin layer is 20% (1.0 mil, 25 μ m) of the film gauge and consists of a blend of Eastar™ 6763 PETG (Eastman Chemical Company, 1.27 g/cc, 0.70 inherent viscosity). index) and 40% LDPE 6811 (The Dow Chemical Company, 0.922 g/cc, 1.2 g/10 minute melt PETG copolyester layer is 10% (0.5 mil, 12 μ m) of the film gauge and consists of 100% A coextruded 5-layer barrier film is made on a conventional upward blown 5-layer The

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innermost layer is extruded at 340°F (171°C). The 5 inch (12.7 cm) diameter tubular blown are extruded at 325°F (163°C); the PETG layer at 400°F (204°C) and the ULDPE-LDPE tube is collapsed to form a 15 inch (38 cm) wide lay flat, slit into two separate webs and wound into film rolls. The Affinity™ layer is extruded at a temperature of 280°F (138°C); the EVA layers is heated at 340°F (171°C). After cooling the blown tube with a dual lip air ring, the

with a fringed surface microstructure. The fringe length is about 300 •m. the film is subjected to a fringing process, wherein the outermost Affinity™ layer is provided In a separate operation and using the equipment described in Example 1 and Fig. 2

the second ply around the perimeter of the pouch die. Films are tested for machine with the heat seal causing the ULDPE-LDPE layer of one ply to bond to the same layer of a heated press (Sentinel 808 heat sealer) using a temperature of 325°F (163°C) and 4 F-1249 using a Mocon Permatran W-600 permeability tester. permeability tester and water vapor transmission rate (WVTR) is tested according to ASTM transmission rate (O₂TR) is tested according to ASTM D-3985 using a Mocon Ox-Tran 1050 both before fringing (initial base film) and after fringing (fringed film). In addition, oxygen direction (MD) and transverse direction (TD) physical properties according to ASTM D-882 each ply is on the outside of the pouch and the ULDPE-LDPE blend layer is on the inside second dwell. Two plies of the film are oriented such that the fringed AffinityTM resin layer of ended pouches of the fringed film are fabricated using a heated 1-liter bag die mounted on The resulting fringed film exhibits a soft, velour or fabric-type feel. Closed

in the centermost layer of the 5 layer asymmetrical structure. The resulting film is subjected 3180 EVA (DuPont, 28% vinyl acetate, 0.95 g/cc, 25 g/10 MFI) is extruded at 310°F (154°C) the modification that the 10% PETG core barrier layer is replaced with a 10% layer of to fringing as described above. A 4-layer coextruded fringed film similar to that described above is produced, with A blend of 96% Saran™ 469 PVDC (The Dow Chemical Company) and 4% Elvax™

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Example 6: Coefficient of Friction (COF)

described in Example 1. Fringe height (H) and Hollowness Index (•) are determined via films are three-layer films and have the same structure and composition of the film 1A microns (6B) and a Hollowness Index (•) of 43, and a fringed film (6C) having a fringe height following instrument conditions: transmitted light using Nomarski interference contrast at optical microscopy using an Olympus Vanox-S Model AH-2 research microscope and the (H) of 95 microns and a Hollowness Index (*) of 60 are tested for their frictional behavior. All portion of the film with a diamond knife on a Leica Ultracut-UCT equipped with an FCS prepared as follows: cross sections having a thickness of 2.5 microns are taken from a magnifications of 100X and 200X with a Polaroid DMC le digital camera. The samples are A smooth base film (6A) (control), a fringed film (6B) having a fringe height (H) of 70

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glass slides containing a suitable immersion oil and analyzed. medium manufactured by Oxford Labware). Embedding consists of placing a few drops of cryosectioning chamber. Films having short fringes (< 75 microns in length) are cut at at 55°C and 30 in. Hg. for approximately 30 minutes. melted wax on the film surface with the fringes. The samples are placed in a vacuum oven Films having longer fringes are embedded in ParaplastTM wax (a tissue embedding The cross sections are placed on

different surfaces: The kinetic coefficient of friction according to ASTM method D1894 is measured for

- a) Fringed side of film (or corresponding side for film 6A) to metal
- b) Fringed Side of Film to Fringed Side of Film

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c) Wet Fringed Side of Film to Wet Metal.

1-3 mm deep. ambient temperature for 24 hours prior to testing. The metal surface is covered with water, For test c), the ASTM method is slightly modified. The film is soaked in deionized water at

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increase over the distance traveled by the test (range tested: 0 to 400 inches). and 6C are 0.1, 0.1 and 1.1, respectively. The COF values for films 6B and 6C are found to obtain increased COF versus the base. The average COF values measured for films 6A, 6B results of test c) (wet film vs. wet metal) show that the fringe must be of a certain length to found to increase over the distance traveled by the sled (tested range: 0 to 400 inches). The measured for films 6A, 6B and 6C are 0.2, 0.3 and 1.4, respectively. The COF for film 6C is causes an increased COF as compared to the smooth film. The average COF values measured for film 6A (COF = 0.1). The results of test b) (film vs. film) show that fringing fringed films 6B (COF = 1.2) and 6C (COF = 2.65) is significantly higher than the COF compared to the smooth film. The average COF of 5 measurements measured on the The results of test a) (fringed vs. metal) show that fringing causes an increased COF

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Example 7: Hot Tack Testing

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peel speed. The hot tack tests are conducted on the samples, whereby one inch wide strips of film are cut and the matrix layer is folded over and pressed to itself by the hot tack tester. test is conducted at 40 psi bar pressure, 0.5 sec dwell time, 0.2 sec delay and 200 mm/min delay time and the seal is pulled at the designated peel speed. Hot tack seals are made at The hot tack tester stays closed for the specified dwell time, it releases the jaws for the Hot tack testing is accomplished using a Topwave Hot Tack Tester Model 52D. The